

Programmer's Reference



ANSYS, Inc.
Southpointe
2600 ANSYS Drive
Canonsburg, PA 15317
ansysinfo@ansys.com
http://www.ansys.com
(T) 724-746-3304
(F) 724-514-9494

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Preface

About the Programmer's Reference (p. 1)

The *Programmer's Reference* (p. 1) provides information about the various programming interfaces available to customers. This manual assumes that you have at least a basic knowledge of programming (a working knowledge of FORTRAN would be very helpful). The two part manual includes:

Part I - Guide to Interfacing with ANSYS

This guide describes a group of utilities as well as a set of FORTRAN routines that you can use to directly access the ANSYS database. You can also use these capabilities to access data in any of the binary files that Mechanical APDL writes or uses.

Part II - Guide to User-Programmable Features

Mechanical APDL provides a set of FORTRAN functions and routines that are available to extend or modify the program's capabilities. Using these routines requires relinking the Mechanical APDL program, resulting in a custom version of Mechanical APDL. The program provides an external commands capability which you can use to create shared libraries available to Mechanical APDL (either from ANSI standard C or FORTRAN). You can use this feature to add custom extensions to Mechanical APDL without the need to rebuild the Mechanical APDL executable.

In addition, you can find the ANSYS Parametric Design Language Guide as part of the Mechanical APDL Help system. This guide was designed for Mechanical APDL users who have some programming skills and wish to leverage the power of the ANSYS Parametric Design Language (APDL) to increase the productivity. APDL is a scripting language that is very similar to FORTRAN. The guide describes how to define parameters (variables), how to create macro programs using APDL, how to use APDL for simple user interaction, how to encrypt an APDL macro, and how to debug an APDL macro.

Note:

The *Programmer's Reference* (p. 1) is offered solely as an aid, and does not undergo the same rigorous verification as the Mechanical APDL product documentation set. Therefore, the *Programmer's Reference* (p. 1) is not considered to be part of the formal program specification as stated in your license agreement.

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Chapter 1: Format of Binary Data Files

Mechanical APDL writes several binary files to store data during an analysis. These files are named <code>Jobname.ext</code>, where <code>Jobname</code> is the name of the analysis that caused the file to be generated and <code>.ext</code> is an extension indicating the type of data in the file.

The following Binary Data File topics are available in this chapter:

- 1.1. Understanding Mechanical APDL Binary Files
- 1.2. Description of the Results File
- 1.3. Description of the Reduced Displacement File
- 1.4. Description of the Reduced Complex Displacement File
- 1.5. Description of the Modal Results File
- 1.6. Description of the Element Matrices File
- 1.7. Description of the Substructure Matrices File
- 1.8. Description of the Component Mode Synthesis Matrices (CMS) File
- 1.9. Description of the Full Stiffness-Mass File
- 1.10. Description of the Substructure Displacement File

1.1. Understanding Mechanical APDL Binary Files

Mechanical APDL-written binary files include the following:

- The following results files, in which Mechanical APDL stores the results of solving finite element analysis problems:
 - Jobname . RST A structural or coupled-field analysis
 - Jobname . RTH A thermal analysis
 - Jobname . RMG A magnetic analysis
- The Jobname . MODE file, storing data related to a modal analysis
- The Jobname . RDSP file, storing data related to a mode-superposition transient analysis.
- The Jobname . RFRQ file, storing data related to a mode-superposition harmonic analysis
- The Jobname . EMAT file, storing data related to element matrices
- The Jobname . SUB file, storing data related to substructure matrices
- The Jobname . FULL file, storing the full stiffness-mass matrix

• The Jobname . DSUB file, storing displacements related to substructure matrices

The files listed above cover almost all users' needs, although there are others. For more information, see the *Basic Analysis Guide*.

1.1.1. Conventions Used to Describe Binary Files

In the information describing the binary file formats:

- Record ID is the identifier for this record. Not all records will have identifiers; they're indicated only for records whose record pointers are stored in a header.
- Type indicates what kind of information this record stores.
- Number of records indicates how many records of this description are found here.
- Record length indicates the number of items stored in the record.

In some record descriptions, actual variable names used may appear in the record contents area.

1.1.2. The Standard Header for Mechanical APDL Binary Files

Each of the Mechanical APDL program's binary files contains a standard, 100-integer file header that describes the file contents. The header contains the items listed below, always in the order shown:

1. 4	T. C
ltem 1	The file number
Item 2	The file format. This item has a value of 1 if the file is small format, -1 if large format.
Item 3	The time, in compact form (that is, 130619 is 13:06:19)
ltem 4	The date, in compact form (that is, 20041023 is 10/23/2004)
Item 5	The units of measurement used. The value of this item is as follows:
	0 for user-defined units
	• 1 for SI units
	• 2 for CSG units
	3 for U. S. Customary units (feet)
	4 for U. S. Customary units (inches)
	5 for MKS units
	6 for MPA units
	• 7 for µMKS units
Item 10	The Mechanical APDL release level in integer form ("X.X" in character form)
Item 11	The date of the Mechanical APDL release
Items 12-14	The machine identifier in integer form (three four-character strings)

The <i>Jobname</i> in integer form (two four-character strings)
The Mechanical APDL product name in integer form (two four-character strings)
The Mechanical APDL special-version label in integer form (one four-character string)
The user name in integer form (three four-character strings)
The machine identifier in integer form (three four-character strings)
The system record size
The maximum file length
The maximum record number
The number of cores used with shared-memory parallel
The number of cores used with distributed-memory parallel
The <i>Jobname</i> (eight four-character strings)
The main analysis title in integer form (20 four-character strings)
The first subtitle in integer form (20 four-character strings)
File compression level
File sparsification key
The split point of the file (0 means the file will not split)
LONGINT of the maximum file length

1.2. Description of the Results File

The next few pages describe the format of the Mechanical APDL results file. (In the following tables, records with a record ID containing an asterisk (*) are those you can read and store into the ANSYS database via the **LDREAD** command.)

Note: The pointers in the solution data headers are relative, not absolute pointers. For example, the 12th item in the solution data header will be relative to a position in the Data Set Index (ptrESL = DSI(i) + ptrESL).

This section explains the contents of the results file; that is, those files with the following extensions:

- .rmg
- .rst
- .rth
- .lnn

The ***XPL** command enables you to explore the contents of certain ANSYS binary files, including the results file. For more information, see Appendix B: Using APDL to List File Structure and Content in the ANSYS Parametric Design Language Guide.

1.2.1. Nomenclature

A load case contains the results for an instance in an analysis. A load case is defined by a load step number and a substep number. A load case is also categorized by a cumulative iteration number and time (or frequency) values. A load case is identified by all three methods in the results file.

The results file does not have to contain all the load cases of an analysis.

A data set is used in this chapter to designate a load case.

For a complex analysis, there will be two data sets for each load case. The first data set contain the real solution and the second contains the imaginary solution.

1.2.2. Standard ANSYS File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 12.

1.2.3. Results File Format

```
*comdeck.fdresu
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc
      ****** description of results file *******
С
      --- used for the following files:
C
С
        .rmq
С
        .rst
       .rth
C
       .lnn(lxx)
      character*8 RSTNM
      parameter (RSTNM='rst
                    resufpL, adrZipL, resuRfpL
     LONGINT
     integer
                    resubk, resuut, resuRbk, resuRut
     common /fdresu/ resufpL, adrZipL, resubk, resuut,
                      resuRfpL, resuRbk, resuRut
   ****** common variable descriptions ********
С
co resufpL
               file position on file resu
                block number for file resu (usually 6)
co resubk
co resuut
                file unit for file resu
                                            (0 if not open) FUN12
c0
               variables for remote modal RST file
    See fddesc for documentation of how binary files are stored
C
      ****** file format *******
С
         recid tells the identifier for this record. Not all records will have
С
             identifiers -- they are only indicated for those records whose
С
             record pointers are stored in a header.
C
         type tells what kind of information is stored in this record:
С
              i - integer
C
С
             dp - double precision
             cmp - complex
C
С
        nrec tells how many records of this description are found here
        lrec tells how long the records are (how many items are stored)
```

```
c recid
           type
                    nrec
                             lrec
                                       contents
                                       standard ANSYS file header (see binhed8 for
                     1
                             100
                                       details of header contents)
C
             i
                     1
                              80
                                       .RST FILE HEADER
C
                                                           nnod, resmax,
С
                                        fun12,
                                                   maxn,
                                                                                numdof,
                                                 nelm,
                                                             kan, nsets, ptrend,
                                        maxe,
                                                                                            (10)
C
C
                                      ptrDSIl, ptrTIMl, ptrLSPl, ptrELMl, ptrNODl,
                                      ptrGEO, ptrCYCl, CMSflg, csEls, units, nSector, csCord, ptrEnd8, ptrEnd8, fsiflag, 0, noffst, eoffst, nTrans, ptrTRAN1, PrecKey, csNds, cpxrst, extopt, nlgeom,
                                                                                           (20)
C
С
C
                                      PrecKey, csNds,
C
                                    AvailDatal, mmass, kPerturb, XfemKey, rstsprs,
С
                                      ptrDSIh, ptrTIMh, ptrLSPh, ptrCYCh, ptrELMh,
С
                                     ptrNODh, ptrGEOh, ptrTRANh, Glbnnod, ptrGNODl, ptrGNODh, qrDmpKy, MSUPkey, PSDkey,cycMSUPkey,
С
                                                                                            (50)
С
С
                             XfemCrkPropTech,cycNoDup,decompMth, nProcSol, mpiWrld,
                                       AMtype, udfrqkey, ptrUDFRQl, ptrUDFRQh, cpxeng,
C
                                                      0,
                                                                 0,
                                                                          0,
                                   AvailDatah,
                                                                                      0,
                                                                                            (70)
C
                                                      0,
                                                                 0,
                                                                            0,
                                            0.
                                                                                       0.
C
                                            0.
                                                      0,
                                                                 0,
                                                                            0,
                                                                                            (80)
С
                                      each item in header is described below:
C
С
                                       fun12 - unit number (resu file is 12)
                                       maxn
                                               - maximum node number of the model
C
                                              - the actual number of nodes used in
С
                                                the solution phase
C
                                       resmax - the maximum number of data sets
С
                                                allowed on the file (defaults to
C
                                                 10000; minimum allowed is 10)
C
                                       numdof - number of DOFs per node
С
С
                                       maxe
                                              - maximum element number of the
                                                finite element model
C
С
                                             - number of finite elements
С
                                       kan
                                              - analysis type
C
                                       nsets - number of data sets on the file
С
                                       ptrend - pointer to the end of the file
                                                  (see ptrEnd8 in 23,24)
С
                                    ptrDSIl,h - 64 bit pointer to the data steps
С
                                                 index table
C
C
                                    ptrTIMl,h - 64 bit pointer to the table of time
С
                                                 values for a load step
                                    ptrLSPl,h - 64 bit pointer to the table of load
С
                                                 step, substep, and cumulative
C
С
                                                 iteration numbers
                                    ptrELMl,h - 64 bit pointer to the element equivalence
C
                                                 table (used when the mesh does not
C
                                                 change during solution)
С
                                    ptrNODl,h - 64 bit pointer to the nodal equivalence
С
С
                                                 table (used when the mesh does not
                                                 change during solution)
C
                                    ptrGEOl,h - 64 bit pointer to the beginning of
C
С
                                                 geometry information (used when the
С
                                                 mesh does not change during solution)
                                    ptrCYCl,h - 64 bit pointer to the table of cyc sym
C
                                                nodal-diameters at each load step
C
                                       CMSflg - CMS results flag: 0-non cms, >0-cms
С
                                       csEls - Cyclic sym # eles in master sector
C
С
                                       units - unit system used
                                                 =-1 - no /UNITS specification
С
                                                 = 0 - user defined units
C
                                                 = 1 - SI
C
                                                 = 2 - CSG
C
                                                 = 3 - U.S. Customary, using feet
С
                                                 = 4 - U.S. Customary, using inches
С
                                                 = 5 - MKS
C
                                                 = 6 - MPA
С
                                                 = 7 - uMKS
```

```
С
                                   nSector - number of sectors for cyclic sym
                                   csCord - Cyclic symmetry coordinate system
С
                                   ptrEnd8 - 64 bit file pointer to the end of
C
С
                                             the file (i.e., length of file)
С
                                   fsiflag - FSI analyis flag
                                   noffst - node offset used in writing file
C
                                   eoffst - elem offset used in writing file
С
                                   nTrans - number of SE transformation vects
С
                                ptrTRAN1,h - 64 bit pointer to SE transformation vects
C
C
                                   PrecKey - 0, double precision
                                             1, single for element results only
C
С
                                             2, single for all data
C
                                   csNds
                                           - Cyclic sym # nds in master sector
                                   cpxrst - complex results flag (0-no, 1-yes)
C
                                   extopt - mode extraction option
С
С
                                   nlgeom - NLGEOM key
                             AvailDatal,h - bits indicating available data any
С
                                             where on the file; see resucm.inc
С
                                          - number of missing mass resp. present
С
                                  kPerturb - key for Linear Perturbation results
С
                                   XfemKey - XFEM flag (set equal to Active_XfenId)
C
                                                       (0=Inactive, 1=Active)
C
                                   rstsprs - bitmask for suppressed items
C
                                   Glbnnod - global number of nodes actually used
C
                                             in the solution phase (== nnod unless
C
С
                                             using Distributed Ansys)
                                ptrGNODl,h - 64 bit pointer to the global nodal
С
C
                                             equivalence table (only used with
                                             Distributed ANSYS and when the mesh
С
                                             does not change during solution)
С
                                   grDmpKy - QR damped calculations key
С
                                   MSUPkey - MSUP results expanded with MSUPCombineModes
C
                                   PSDkey - PSD key
C
                                cycMSUPkey - rst file format is for subsequent cyclic MSUP
C
С
                                             (only base results on file)
                           XfemCrkPropTech - XFEM crack propagation technology
C
С
                                       (0=phantom node-based, 1=singularity-based)
С
                                  cycNoDup - no duplicate sector created
                                 decompMth - domain decomposition method employed with DMP
C
С
                                             (>= 0 = MESH; -1 = FREQ, -2 = CYCHI)
                                  nProcSol - when the domain decomposition method for DMP
С
                                             is FREQ or CYCHI --> number of processes used for
C
                                             each frequency or cyclic harmonic index solution
C
                                   \ensuremath{\mathsf{mpiWrld}} - unique identifier for determining which results
C
С
                                             file belongs to
                                    AMtype - additive manfacturing (AM) flag
C
                                  udfrqkey - Key for format and writing of undamped frequency record
C
С
                       ptrUDFRQl,ptrUDFRQh - pointer to undamped frequency record
C
                                    cpxeng - 0 = default; lrec = 11
                                             1 = element ENG record contains additional quantities;
C
                                                 lrec = 21
С
С
С
                                    Note: ptrXXX are relative to beginning of file
           i
                           numdof
                                    Degrees of freedom per node
C
C
                                    DOF reference numbers are:
          UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8
С
          AZ = 9, VX = 10, VY = 11, VZ = 12, GFV1 = 13, GFV2 = 14, GFV3 = 15, WARP = 16
C
          CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
С
С
          EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
          С
С
          TE17=49, TE18=50, TE19=51, TE20=52, TE21=53, TE22=54, TE23=55, TE24=56
С
          TE25=57, TE26=58, TE27=59, TE28=60, TE29=61, TE30=62, TE31=63, TTOP=64
C
                                     (curdof(i), i=1, numdof)
С
С
    NOD
                           nnod
                                    Nodal equivalence table. This table equates
С
                                    the number used for storage to the actual
                                    node number
С
C
                                     (Back(i), i=1, nnod)
```

0 0 0 0	ELM	i	1	nelm	Element equivalence table. The ANSYS program stores all element data in the numerical order that the SOLUTION processor solves the elements. This table equates the order number used to the actual element number
0 0 0 0	GNOD	i	1	Glbnnod	Global nodal equivalence table. This table equates the number used for storage to the actual node number. Only written by the master process in Distributed Ansys (GlbBack(i),i=1,Glbnnod)
	DSI	i	1	2*resmax	Data sets index table. This record contains the record pointers for the beginning of each data set. The first resmax records are the first 32 bits of the index, the second resmax records are the second 32 bits. To create the 64 bit pointer, use: LONGPTR = largeIntGet (first, second) Read the solution data header as follows: call bioBasePut (nblk,LONGPTR) loc = bioiqr (nblk,12) call biord (nblk,loc, The rest of the file reading continues to use the ptrXXX's that are in the headers.
с с	TIM	dp	1	resmax	Time/freq table. This record contains the time (or frequency) values for each data set.
0 0 0 0	UDFRQ	dp	1	resmax	Undamped freq table. This record contains the undamped frequency values for each data set in a QRDAMPED modal analysis. This record exists only when qrDmpKy = 1 (see the results file header for qrDmpKy)
с с	LSP	i	1	3*resmax	Data set identifiers. This record contains the load step, substep, and cumulative iteration numbers for each data set.
С	CYC	i	1	resmax	Cyclic symmetry harmonic index
С	TRAN	dp	nTran	25	Substructure transformation vectors
С	GEO	i	1	80	Geometry data header
				q	0, maxety, maxrl, nnod, nelm, maxcsy, ptrETY, ptrREL, ptrLOC, ptrCSY, (10) ptrEID, maxsec, secsiz, nummat, matsiz, ptrMAS, csysiz, elmsiz, etysiz, rlsiz, (20) ptrETYl, ptrETYh, ptrRELl, ptrRELh, ptrCSYl, ptrCSYh, ptrLOCl, ptrLOCh, ptrEIDl, ptrEIDh, (30) ptrMASl, ptrMASh, ptrSECl, ptrSECh, ptrMATl, ptrMATh, ptrCNTl, ptrCNTh, ptrNODl, ptrNODh, (40) ptrELMl, ptrELMh, Glblenb,ptrGNODl,ptrGNODh, maxn,NodesUpd, lenbac, numcomp, mxcmpsz, (50) trCOMPl,ptrCOMPh,nMatProp, nStage, maxMSsz, ptrMSl, ptrMSh, nCycP,ptrCycPl,ptrCycPh, (60) numety, numrl, numcsy, numsec, mapFlag, cysCSID, 0, 0, 0, 0, (70) 0, 0 0, 0, 0, 0, 0, 0 0, 0, 0, 0, 0 0, 0, 0, 0, 0 0, 0, 0, 0, 0 0, 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0, 0 0, 0 0, 0, 0 0, 0 0, 0, 0 0,
C					<pre>maxrl - the maximum real constant reference number in the model</pre>
C					nnod - the number of defined nodes in the

```
С
                                               model
                                             - the number of defined elements in
С
                                      nelm
                                               the model
C
                                      maxcsy - the maximum coordinate system
С
С
                                               reference number in the model
                                      ptrETY - pointer to the element type index
C
С
                                               table
                                      ptrREL - pointer to the real constant
С
                                               index table
C
                                      ptrLOC - pointer to the nodal point
                                               locations
C
C
                                      ptrCSY - pointer to the local coordinate
C
                                               system index table
                                      ptrEID - pointer to the element index
C
C
                                               table
                                      maxsec - the maximum section
C
                                               reference number in the model
C
                                      secsiz - the maximum size that any
С
С
                                               section record may have
                                      nummat - the number of materials
C
                                               in the model
C
                                      matsiz - the maximum size that any material
C
                                               property or table may have
C
                                      ptrMAS - pointer to the diagonal mass matrix
C
                                      csysiz - the number of items describing a
C
С
                                               local coordinate system (usually
С
                                               24)
                                      elmsiz - the maximum number of nodes that a
C
С
                                               defined element may have
                                      etysiz - the number of items describing an
C
                                               element type(=IELCSZ from echprm.inc)
C
                                      rlsiz - the maximum number of items
C
C
                                               defining a real constant (0, if no
                                               real constants are defined)
C
                                   ptrETYl,h - 64 bit pointer to element type data
C
                                   ptrRELl,h - 64 bit pointer to real constant data
C
C
                                   ptrCSYl,h - 64 bit pointer to coordinate system data
                                   ptrLOCl,h - 64 bit pointer to nodal locations
C
                                   ptrEIDl,h - 64 bit pointer to element data ptrSECl,h - 64 bit pointer to section data
C
C
                                   ptrMAT1,h - 64 bit pointer to material data
С
                                   ptrCNTl,h - 64 bit pointer to element centroids
C
                                   ptrNODl,h - 64 bit pointer to nodal equivalence table
C
C
                                   ptrELMl,h - 64 bit pointer to element equivalence table
                                     Glblenb - global number of nodes actually used
C
                                               in the solution phase (== lenbac unless
C
                                               using Distributed Ansys)
C
С
                                   ptrGNOD1,h- 64 bit pointer to the global nodal
C
                                               equivalence table (only used with
                                               Distributed ANSYS and when the mesh
C
                                               does not change during solution)
С
С
                                      maxn
                                            - maximum node number of the model
                                    NodesUpd - 1, node coords have been updated
C
                                      lenbac - the actual number of nodes used in
C
                                               the solution phase
C
                                     numcomp - number of components/assemblies stored
C
                                               (only node/elem components/assemblies)
C
                                     mxcmpsz - maximum size (in integer words) that any
C
                                               component/assembly record may have
C
                                  ptrCOMP1,h - 64 bit pointer to component/assembly data
С
                                    nMatProp - number of properties stored per material
C
                                     nStage - number of stages
maxMSsz - maximum size (in integer words) that a
С
С
                                               stage record can have
C
                                    ptrMSl,h - 64 bit pointer to multistage (MS) cyclic
                                               analysis data
C
С
                                       nCycP - number of cyclic edge node pair tables (CYCLIC and MS)
C
                         ptrCycPl, ptrCycPh - pointers to cyclic edge node pair tables
                                      numety - the number of defined element types
C
                                               in the model
С
                                      numrl - the number of defined real
```

```
constants in the model
С
                                     numcsy - the number of defined coordinate
С
                                               systems in the model
С
                                     numsec - the number of defined sections in
С
С
                                               the model
                                    mapFlag - flag to indicate format of mapping
C
                                               index vectors for element types,
С
                                               real constants, coordinate systems,
С
                                               and sections.
C
                                               = 0, old format with 1 vector of
С
                                                    maxDef len
C
С
                                               = 1, new format with 2 vectors
С
                                                    each having numDef len
                                    cycCSID - coordinate system number (CYCLIC and MS)
C
                                               = 0 is ignored (must be cylindrical)
С
С
                                     Note: ptrXXX are relative to beginning of file
С
                                     if mapFlag == 0 --> Length = maxety
С
    ETY
             i
                     1
                            varies
                                     The element types index table. This record
С
С
                                     contains record pointers for each element
                                     type description.
С
С
                                    (Relative to ptrETYPL)
С
                                     if mapFlag == 1 --> Length = numety
C
                                     Elemnt type external number mapping.
С
                                     This record maps the number used for storage
С
C
                                     to the actual element type number
             i
                            numety
                                     Only exists if mapFlag ==1
C
                                     The element types index table. This record
C
С
                                     contains record pointers for each element
                                     type description.
C
С
                                    (Relative to ptrETYPL)
                                     Element type description. Each of these
             i
                   numety etysiz
C
С
                                     records is pointed to by a record pointer
С
                                     given in the record labeled ETY. See
                                     routines echprm and elccmt for a complete
С
                                     description of the items stored here.
C
                                     These items are typically stored into the
С
                                     IELC array, and are used to determine the
C
С
                                     element type characteristics at runtime.
                                     The following items are typically of
С
С
                                     interest:
                                     * Item 1
                                                   - element type reference number
С
С
                                     * Item 2
                                                   - element routine number
                                     * Items 3-14 - element type option keys
С
С
                                                     (keyopts)
С
                                                   - DOF/node for this element
                                     * Item 34
                                                     type. This is a bit mapping
С
                                                     of the DOF/node.
С
                                     * Item 61
                                                   - number of nodes for this
С
C
                                                     element type (nodelm)
                                                   - number of nodes per element
С
                                      * Item 63
С
                                                     having nodal forces, etc.
                                                     (nodfor)
С
                                                   - number of nodes per element
С
                                     * Item 94
С
                                                     having nodal stresses, etc.
                                                     (nodstr). This number is the
C
                                                     number of corner nodes for
С
                                                     higher-ordered elements.
С
                                                     NOTE- the /config NST1 option
C
С
                                                     may suppress all but one node
                                                     output of nodal ENS EEL EPL
С
                                                     ECR ETH and ENL for any given
С
С
                                                     element.
                                     if mapFlag == 0 --> Length = maxrl
C
                            varies
                                     Real constants index table. The record
C
```

C C					contains record pointers for each real constant set. (Relative to ptrRELL)
0 0 0					if mapFlag == 1> Length = numrl Real constants external number mapping. This record maps the number used for storage to the actual real constant number
c c c c c		i	1	numrl	Only exists if mapFlag ==1 Real constants index table. The record contains record pointers for each real constant set. (Relative to ptrRELL)
		dр	numrl	varies	Element real constant data. These records contain real constant data used for the elements. (See the ANSYS Elements Reference manual for values for a specific element.) Each of these records is pointed to by a record pointer given in the record labeled REL. The length of these records varies for each element type (actual length is returned from routine BINRD8).
	CSY	i	1	varies	if mapFlag == 0> Length = maxcsy Coordinate systems index table. This record contains the record pointers for each coordinate system set. The ANSYS program writes coordinate systems only if local coordinate systems were defined. If a local system was defined, the predefined global systems 1 to 2 also will be written. The global Cartesian system 0 will never be written. (Relative to ptrCSYSL) if mapFlag == 1> Length = numcsy Coordinate system external number mapping. This record maps the number used for storage
		i	1	numcsy	Only exists if mapFlag ==1 Coordinate systems index table. This record contains the record pointers for each coordinate system set. The ANSYS program writes coordinate systems only if local coordinate systems were defined. If a local system was defined, the predefined global systems 1 to 2 also will be written. The global Cartesian system 0 will never be written. (Relative to ptrCSYSL)
		dp	numcsy	csysiz	Coordinate system description. These records contain coordinate system data for each coordinate system defined. Each of these records is pointed to by a record pointer given in the record labeled SYS. The items stored in each record: * Items 1-9 are the transformation matrix. * Items 10-12 are the coordinate system origin (XC,YC,ZC). * Items 13-14 are the coordinate system parameters (PAR1, PAR2). * Items 16-18 are the angles used to define the coordinate system. * Items 19-20 are theta and phi singularity keys.

```
* Item 21 is the coordinate system type
С
                                       (0, 1, 2, or 3).
С
                                      * Item 22 is the coordinate system reference
C
С
                                       number.
                              7
                                     Node, X, Y, Z, THXY, THYZ, THZX for each node
    LOC
C
            dρ
                   nnod
                                     Nodes are in node number order
С
    EID
             i
                                     Element descriptions index table. This
C
                     1
                           2*nelm
C
                                     record contains the record pointers for each
                                     element description. The first nelm values
С
                                     are the lower 32 bits and the second nelm
С
                                     values are the higher 32 bits. The order
С
                                     of the elements is the same as the order
C
                                     in the element equivalence table.
С
                                    (Relative to ptrEIDL)
С
                   nelm 10+nodelm Element descriptions. Each of these records
С
С
                                     is pointed to by a record pointer given in
                                     the record labeled EID. The length of these
С
                                     records varies for each element (actual
C
                                     length is returned from routine BINRD8).
С
С
                                     nodelm shown here is the number of nodes for
                                     this element. Its value is defined in the
С
                                     element type description record.
C
С
                                     The items stored in each record:
C
                                        mat, type, real, secnum, esys,
                                      death, solidm, shape, elnum, baseeid,
С
                                      nodes
С
                                    each item is described below:
C
                                             - material reference number
С
                                      mat
                                             - element type number
С
                                      type
                                             - real constant reference number
                                      real
С
С
                                      secnum - section number
С
                                      esys
                                             - element coordinate system
                                      death - death flag
С
С
                                                = 0 - alive
                                                = 1 - dead
С
                                      solidm - solid model reference
С
                                      shape - coded shape key
C
                                      elnum - element number
С
                                      baseeid- base element number
С
С
                    (applicable to reinforcing elements only)
                                      nodes - node numbers defining the element
С
С
                                                (See the ANSYS Elements Reference
С
                                               for nodal order of an element)
                   nelm
                                     Centroid record for each element
С
    CENT
            dр
    MAS
            dp
                     1 nnod*numdof Diagonal mass matrix
C
                                     if mapFlag == 0 --> Length = maxsec
    SEC
             i
                     1
                            varies
C
                                     Section index table. The record
С
                                     contains record pointers for each
С
                                     section set.
C
С
С
                                     if mapFlag == 1 --> Length = numsec
                                     Section external number mapping.
C
С
                                     This record maps the number used for storage
                                     to the actual section number
С
С
             i
                     1
                            numsec
                                     Only exists if mapFlag ==1
                                     Section index table. The record
С
                                     contains record pointers for each
С
C
                                     section set.
                                     Element section data. These records
C
            dр
                   numsec varies
                                     contain section data used for the
```

c c c c		elements. Each of these records is pointed to by a record pointer given in the record labeled SEC. The length of these records varies for each section type (actual length is returned from routine BINRD8).
C MAT C C C C C C C C C C	i 1 3+nummat*(nMa	<pre>Total Sz = Header Data + (nMatProp+1)*Number of Materials + Tail The 1st 3 integers contain the header information 1) Version number (-101 for differentiation from prev rst data) 2) Header size which is 3 3) Size of the index array Ith material ID is stored at Data(3+((I-1)*(nMatProp+1))+1)</pre>
c c c c c c		Material record pointers for the material ID at Ith position is stored from location 3+(I-1)*(nMatProp+1)+2 to 3+(I-1)*(nMatProp+1)+1+nMatProp The record includes MP pointers followed by TB pointers for a single Material ID Last 2 data contains the active table number and material ID
c c c c c c c	dp nummat varies	Material property data. These records contain material property data used for the elements. Each of these records is pointed to by a record pointer given in the record labeled MAT. The length of these records varies for each material property type (actual length is returned from routine BINRD8).
C NOD C C	i 1 lenbac	Nodal equivalence table. This table equates the number used for storage to the actual node number (Back(i),i=1,lenbac)
C ELM C C C	i 1 nelm	Element equivalence table. The ANSYS program stores all element data in the numerical order that the SOLUTION processor solves the elements. This table equates the order number used to the actual element number
C GNOD C C C	i 1 Glblenb	Global nodal equivalence table. This table equates the number used for storage to the actual node number. Only written by the master process in Distributed Ansys (GlbBack(i),i=1,Glblenb)
C COMP C C C C C C C C C C	i numCmp varies	Component/assembly data. The first word will describe the type of component or assembly (1=node component, 2=elem component, 11->15 assembly) For components, values 2->9 are the component name converted to integers. The remaining values are the list of nodes/elems in the component. For assemblies, the remaining values (in groups of 8) are the component names converted to integers.
C MS C C C C C C	nStage i PARMSIZEW+25	1-PARMSIZEW packed stage name PARMSIZEW+1 SEkey, Nsector, Hindex, Numoff cyclic CE min/max downstream interstage CE min/max upstream interstage CE min/max number of nodes, min/max node numbers (base only)
с с	dp 10	number of elements, min/max element numbers (base onl Stage dp data. 1-6 min/max nodal coordinates of the stage

```
1 nStage*nStage Table of connection between stages
C
    CYCP
                            varies
                                      Record exists only if the number of tables nCycP and
C
С
                                      associated pointers are non-zero.
С
                                      nCycP = 1 for CYCLIC
                                            = number of non-superelement and non-3D stages (MS)
C
                                      Maximum record size = number of nodes defined
С
                                      Record content:
С
                                         1:nCycP
                                                  = number of nodes for each table (iCycP)
C
С
                                         nCycP+1... = node pairs of each table
                                      Effective record size = nCycP + sum(iCycP*2)
C
C
         The solution information is stored starting at this point in the file.
         The remaining records on the file are repeated as a group nsets times
C
         (once for each data set). Item nsets is defined in the file header.
C
         Each set of data is pointed to by a record pointer given in the record
C
         labeled DSI.
С
                              200
                                       Solution data header.
С
                                                         nnod,
                                                                           itime.
                                          0.
                                                nelm.
                                                                maskl,
С
                                       iter, ncumit,
                                                         nrf, cs_LSC,
                                                                           nmast,
                                                                                    (10)
С
                                                        ptrRF,
                                     ptrNSL, ptrESL,
                                                                 ptrMST,
                                                                            ptrBC,
С
                                                         isym, kcmplx, numdof,
                                                mode,
C
                                     rxtrap,
                                                                                    (20)
                                                DOFS,
                                                         DOFS,
                                                                   DOFS,
С
                                       DOFS.
                                                DOFS,
                                                         DOFS,
                                                                   DOFS,
                                       DOFS,
                                                                             DOFS,
С
                                                                                    (30)
C
                                       DOFS,
                                                DOFS,
                                                         DOFS,
                                                                   DOFS,
                                                                             DOFS,
                                                DOFS,
                                                         DOFS,
                                                                   DOFS,
                                                                             DOFS,
С
                                       DOFS,
                                                DOFS,
                                                         DOFS,
                                                                             DOFS,
С
                                      DOFS.
                                                                   DOFS,
                                      DOFS,
                                               DOFS,
                                                         DOFS,
                                                                  DOFS,
                                                                            DOFS,
С
                                                                                    (50)
С
                                      title,
                                               title,
                                                        title,
                                                                  title,
                                                                            title,
                                               title,
                                                        title,
                                                                  title,
                                                                            title,
C
                                      title,
                                                                                    (60)
С
                                      title,
                                               title,
                                                        title,
                                                                  title,
                                                                            title,
С
                                     title,
                                               title,
                                                        title,
                                                                 title,
                                                                            title,
                                                                                    (70)
                                     stitle, stitle, stitle, stitle,
С
С
                                     stitle, stitle, stitle, stitle,
                                     stitle, stitle, stitle, stitle,
С
                                     stitle, stitle, stitle, stitle, stitle, dbmtim, dbmdat, dbfncl, soltim, soldat,
С
                                                                                    (90)
С
                                                                   0, ptrEXT,
                                     ptrOND, ptrOEL,
                                                         0,
С
                                                                                    (100)
                                          0,
                                                   0, ptrEXTl, ptrEXTh, ptrNSLl,
С
                                    ptrNSLh, ptrRFl, ptrRFh, ptrMSTl, ptrMSTh,
                                                                                    (110)
C
                                                        0,
                                    ptrBCl, ptrBCh, 0, 0, ptrONDl,
ptrONDh, ptrOELl, ptrOELh, ptrESLl, ptrESLh,
С
С
                                                                                    (120)
С
                                   ptrOSL1, ptrOSLh, sizeDEAD, ptrDEAD1, ptrDEADh,
                                   PrinKey, numvdof, numadof, ptrNARl, ptrNARh,
С
                                                                                    (130)
С
                                   ptrVSLl, ptrVSLh, ptrASLl, ptrASLh, nMSHI,
                                  ptrMSHIl,ptrMSHIh,     0,numRotCmp,     0,
ptrRCMl, ptrRCMh, nNodStr,     0,ptrNDSTRl,
С
                                                                                    (140)
C
                                 ptrNDSTRh, AvailDatal, geomID, ptrGEOl, ptrGEOh,
С
                                      k2d3d, maskh, AMstep, inverse, AvailDatah,
С
                                                  0,
С
                                          0,
                                                           0,
                                                                      0, ptrCINT,
                                                                      0,
                                    lenCINT, ptrXFEM, lenXFEM,
С
                                                                                0,
                                                   0,
                                                             0,
                                          0,
                                                                      0,
                                                                                0,
                                                                                    (170)
C
С
                                          0,
                                                   0,
                                                             0,
                                                                      0,
                                                                                Ο.
С
                                          0,
                                                   0,
                                                             0,
                                                                      0,
                                                                                0,
                                                                                    (180)
                                                   0,
                                                             0,
                                                                      0,
                                          0,
C
                                                                                0.
С
                                          Ο,
                                                   0,
                                                             0,
                                                                      0,
                                                                                0,
                                                                                    (190)
С
                                          0,
                                                   0,
                                                             0,
                                                                      0,
                                                                                0,
                                          0,
                                                   0,
                                                             Ο,
                                                                      0,
                                                                                    (200)
C
                                     each item in header is described below:
С
                                      nelm
                                            - number of elements
C
                                            - number of nodes
                                     nnod
C
                                     maskl,h - bitmask for the existence of
С
С
                                               several records. If a bit is set
                                               here, it indicates that the
C
                                               corresponding record exists on the
С
С
```

```
The items in the bitmask that
С
С
                                                correspond to each record are shown
С
                                                in the record descriptions below.
                                       itime - loadstep
С
С
                                       iter
                                             - iteration number
                                       ncumit - cumulative iteration number
C
                                               - number of reaction forces
С
                                       cs_LSC - cyclic symmetry count of the
С
                                                load step for this SOLVE
C
C
                                       nmast - number of masters
                                       ptrNSL - 32-bit pointer to nodal solution
C
                                      ptrESL - 32-bit pointer to element solution
ptrRF - 32-bit pointer to reaction forces
С
C
                                       ptrMST - 32-bit pointer to the masters
C
                                       ptrBC - 32-bit pointer to the boundary conditions
C
                                       rxtrap - key to extrapolate integration
С
                                                point results to nodes
C
                                                = 0 - move
= 1 - extrapolate unless active
С
С
                                                      non-linear
C
                                                = 2 - extrapolate always
C
                                              - mode number of harmonic loading
                                       mode
C
                                                (for cyclic symmetry: this is cs_LSF
C
                                                = first load step for this SOLVE)
C
                                              - symmetry for harmonic loading
C
                                       isvm
                                                (for cyclic symmetry: this is cs_LSL
С
С
                                                = last load step for this SOLVE)
C
                                       kcmplx - complex key
                                                = 0 - real
= 1 - imaginary
С
C
                                       numdof - number of DOFs/nodes for this data
C
C
                                                set
C
                                       DOFS
                                             - DOF/node reference numbers (numdof
                                                values)
C
                                       title - main title (in integer form)
С
                                      stitle1 - 1st subtitle (in integer form)
C
С
                                       dbmtim - time (in compact form) when the
С
                                                database was last modified
                                       dbmdat - date (in compact form) when the
C
С
                                                database was last modified
                                       dbfncl - number of times that the database
С
                                                was modified
C
                                       soltim - time (in compact form) when the
C
C
                                                solution for this data set was done
                                       soldat - date (in compact form) when the
С
С
                                                solution for this data set was done
                                       ptrOND - 32-bit pointer to the ordered node
C
С
                                                list (load case files only)
                                       ptrOEL - 32-bit pointer to the ordered element
C
                                                list (load case files only)
С
                                       ptrEXT - 32-bit pointer to header extension
С
                                   ptrEXTl,h - 64-bit pointer to header extension
С
                                   ptrNSLl,h - 64-bit pointer to nodal solution
C
                                   ptrRFl,h - 64-bit pointer to reaction forces
C
                                   ptrMSTl,h - 64-bit pointer to the masters
C
                                   ptrBCl,h \, - 64-bit pointer to the boundary conditions ptrONDl,h \, - 64-bit pointer to the ordered node
С
C
                                                list (load case files only)
C
                                   ptrOELl,h - 64-bit pointer to the ordered element
С
С
                                                list (load case files only)
                                   ptrESL1,h - 64-bit pointer to element solution
C
                                   ptrOSLl,h - 64-bit pointer to extra solution vector
С
                                                (to be used later for rezoning project)
С
                                   sizeDEAD - size of dead element list
C
                                  ptrDEADl,h - 64-bit pointer to dead element list
C
                                      PrinKey - principal stress key:
C
С
                                                0, use rstsprs (if bit 26 set, no prin)
С
                                                1, principals are written for this set
                                               -1, no principals are written for this set
C
                                      numvdof - number of velocity dofs
С
                                      numadof - number of acceleration dofs
```

```
ptrNARl,h - 64-bit pointer to nodal averaged
С
С
                                              results solution
                                 ptrVSLl,h - 64-bit pointer to transient velocity
С
С
                                              solution
                                 ptrASLl,h - 64-bit pointer to transient acceleration
С
                                              solution
C
                                            - length of multistage harmoninc index table
С
                                 ptrMSHIl,h- 64-bit pointer to multistage harmonic
С
                                             index table
C
C
                                 numRotCmp - number of rotating components
                                 ptrRCM,h - 64-bit pointer to RCM
C
                                   nNodStr - 0, no nodal component stresses
С
С
                                              1, one set (TOP for shells)
                                              2, two sets (TOP,BOT for shells)
C
                                              3, three sets (TOP,BOT,MID)
С
                               ptrNDSTRl,h - 64 bit pointer to nodal component str
С
                              AvailDatal,h - bits indicating available data
С
                                              in this data set; see resucm.inc
С
                                    geomID - number identifying which geometry (mesh)
С
                                              is used for this set of data (when mesh
С
                                              does not change during solution this should
C
                                              always be equal to 1)
C
С
                                 ptrGEOl,h - 64 bit pointer to geometry data (when mesh
                                              does not change during solution this points
C
                                              to first GEO record near start of file)
C
                                       k2d3d - key indicator 3d run on for 2d 3d analysis
С
                                      AMstep - additive manufacturing process step
С
C
                                               BUILD, COOL, HIP, REMOVE, USER, HEATTREAT
                                                      3 4 5
С
                                                                      6
                                    inverse - 1, the result data is from an inverse analysis
С
                                               0, the result data is from a forward analysis
C
                                               2, the result data is from a forward analysis
C
                                                     after inverse analysis
C
                                    ptrCINT - pointer to CINT record
lenCINT - length of CINT record (doubles)
С
С
                                    ptrXFEM - pointer to XFEM record
С
С
                                    lenXFEM - length of XFEM record (doubles)
С
                                          0 - position not used
С
С
                                    Note: ptrXXX are relative to ptrDSI, except ptrGEO
                                           which is relative to the beginning of the file
С
                     1
                              100
                                    Solution header - double precision data
C
            dp
С
                                      timfrg, lfacto, lfactn, cptime,
                                                                            tref,
С
                                       tunif,
                                                0,
                                                        0,
                                                                     0, dsfpinc,
                                                                                    (10)
                                       accel.
                                                accel,
                                                         accel,
C
                                                                  omega, omega,
С
                                       omega,
                                               omega,
                                                        omega,
                                                                 omega, omegacg,
С
                                     omegacg, omegacg, omegacg, omegacg,
                                      cgcent, cgcent, fatjack, fatjack,
C
                                                                                    (30)
                                       dval1,
                                               dval2,
                                                        dval3,
                                                                   0,
С
                                                                               0,
                                                  0,
                                                           0,
                                          0,
С
                                                                      0,
                                                                               0,
                                                                                    (40)
                                                            0,
C
                                          Ο,
                                                   0,
                                                                     0,
                                                                               0,
                                                   0,
                                                             0,
                                          0,
                                                                      0,
                                                                               0,
C
                                      timdat, timdat, timdat, timdat,
C
                                      timdat, timdat, timdat, timdat,
timdat, timdat, timdat, timdat,
С
                                                                          timdat,
                                                                                    (60)
С
                                                                          timdat,
                                      timdat, timdat, timdat, timdat,
C
                                      timdat, timdat, timdat, timdat,
С
С
                                      timdat, timdat, timdat, timdat,
                                                                                    (80)
                                      timdat, timdat, timdat, timdat,
С
                                      timdat, timdat, timdat, timdat, timdat, timdat, timdat, timdat,
С
С
                                                                          timdat,
                                      timdat, timdat, timdat, timdat, timdat
C
                                                                                    (100)
                                   each item is described below:
C
C
                                      timfrq - time value (or frequency value,
                                                for a modal or harmonic analysis)
C
                                      lfacto - the "old" load factor (used in
С
C
                                                ramping a load between old and new
```

```
С
                                                  values)
                                       lfactn - the "new" load factor
С
                                       cptime - elapsed cpu time (in seconds)
C
                                                - the reference temperature
С
С
                                       tunif
                                               - the uniform temperature
                                       dsfpinc - incident power of diffuse sound field
C
                                               linear acceleration termsangular velocity (first 3 terms) and
С
С
                                       omega
                                                  angular acceleration (second 3 terms)
C
C
                                       omegacg - angular velocity (first 3 terms) and
                                                  angular acceleration (second 3 terms)
C
С
                                                  these velocity/acceleration terms are
С
                                                  computed about the center of gravity
                                       cgcent - (x,y,z) location of center of gravity
C
                                       fatjack - FATJACK ocean wave data (wave height
С
С
                                                  and period)
                                         dvall - FATJACK ocean wave direction
С
                                         dval2 - machine rpm
dval3 - central frequency of tune band
С
С
                                        timdat - load data (slot 53 is substep convergence key)
С
    EXT
             i
                      1
                                200
                                       Header extension
C
                                       positions
                                                  1-32 - current DOF for this
С
С
                                                            result set
                                       positions 33-64 - current DOF labels for
C
С
                                                            this result set
                                       positions 65-84 - The third title, in
С
C
                                                            integer form
                                       positions 85-104 - The fourth title, in
С
С
                                                            integer form
                                       positions 105-124 - The fifth title, in
С
                                                            integer form
C
C
                                       position 125 - unused
                                       position 126 - unused position 127 - numvdof, number of velocity
С
С
                                                       items per node (ANSYS
C
                                                       transient)
С
                                       position 128 - numadof, number of
С
                                                       acceleration items per
С
                                                       node (ANSYS transient)
С
                                       position 131-133 - position of velocity
С
                                                           in DOF record
С
                                                           (ANSYS transient)
C
С
                                       position 134-136 - position of acceleration
С
                                                           in DOF record
С
                                                           (ANSYS transient)
                                       position 137-142 - velocity and
С
С
                                                           acceleration labels
С
                                                           (ANSYS transient)
                                       position 143 - number of stress items
C
                                                       (6 or 11); a -11 indicates
С
С
                                                       to use principals directly
С
                                                       and not recompute (for PSD)
                                       position 144-146 - position of rotational
C
                                                           velocity in DOF record
C
С
                                                           (ANSYS transient)
                                       position 147-149 - position of rotational
С
                                                           accel. in DOF record
C
С
                                                           (ANSYS transient)
                                       position 150-155 - rotational velocity and
С
                                                           acceleration labels
C
С
                                                            (ANSYS transient)
                                       position 160 - pointer to CINT results
С
                                       position 161 - size of CINT record
C
С
                                       position 162 - Pointer to XFEM/SMART-crack surface information
                                       position 163 - size of crk surface records
С
                                       position 164-200 - unused
С
    GEO
                                       Entire geometry record for this set of
C
                            varies
                                       results data. Note, when the mesh does not
С
                                       change during solution, this pointer will
C
```

c c c c			simply point to the original GEO record stored at the start of the file. When the mesh changes the new geoemtry data will be stored here. The geomID can be used to determine when the mesh changes.
c * NSL c c c c c c c c c c c c c c c c c c c	dp	1 nnod*Sumdof	The DOF solution for each node in the nodal coordinate system. The DOF order is the same as shown above in the DOF number reference table. The nodal order is the same order given above in the nodal equivalence table. If a DOF for a node isn't valid, a value of 2.0**100 is used. Note 1: Sumdof = numdof Note 2: If, upon reading of this record, there is less than nnod*Sumdof items in the record, then only a selected set of nodes were output. Another record follows (integer, less than nnod long) which contains the list of nodes for which DOF solutions are available. (bit 10 (PDBN) in mask)
c VSL c c c	dp	1 nnod*numvdof	The velocity solution for each node in the nodal coordinate system. The description for the DOF solution above also applies here. ANSYS transient. (bit 27 (PDVEL) in mask)
C ASL C C C	dp	1 nnod*numadof	The acceleration solution for each node in the nodal coordinate system. The description for the DOF solution above also applies here. ANSYS transient. (bit 28 (PDACC) in mask)
c OSL	dp	1 nnod*Sumdof	extra solution vector for element team rezoning project. To be used later.
C RF C C C	LONG	1 nrf	Reaction force DOFs. This index is calculated as (N-1)*numdof+DOF, where N is the position number of the node in the nodal equivalence table, and DOF is the DOF reference number. (bit 11 (PDBR) in mask)
C * C C	dp	1 nrf	Reaction forces. The force values are ordered according to the DOF order shown above in the DOF number reference table. (bit 11 (PDBR) in mask)
c MST c c c	LONG	1 nmast	Master DOF list. This index is calculated as (N-1)*numdof+DOF, where N is the position number of the node in the nodal equivalence table, and DOF is the DOF reference number.
c BC c c c c c c c c c	i	1 40	Boundary condition index table. (bit 23 (PDBBC) in mask) numdis,ptrDIX,ptrDIS,numfor,ptrFIX, ptrFOR,format, 0,
С		6	each item is described below: numdis - number of nodal constraints
C			ptrDIX - pointer to the table of nodes

					having nodal constraints ptrDIS - pointer to nodal constraint values numfor - number of nodal input force
	DIX	i	1	numdis	if format == 0> Nodal constraint DOF. This index is calculated as N*32+DOF, where N is the node number and DOF is the DOF reference number. Values are in the same order as the DOF number reference table. if format == 1> Nodal constraint node numbers.
0 0 0 0		i	1	numdis	<pre>if format == 0> does not exist. if format == 1> Nodal constraint DOF. Values are in the same order as the DOF number reference table.</pre>
0 0 0	DIS	dp	1	4*numdis	Nodal constraints. This record contains present and previous values (real and imaginary) of the nodal constraints at each DOF.
	FIX	i	1	numfor	if format == 0> Nodal input force DOFs. This index is calculated as N*32+DOF, where N is the node number and DOF is the DOF reference number. Values are in the same order as the DOF number reference table. if format == 1> Nodal input force node numbers.
0 0 0 0		i	1	numfor	<pre>if format == 0> does not exist. if format == 1> Nodal input force DOF. Values are in the same order as the DOF number reference table.</pre>
с с	FOR	dp	1	4*numfor	Nodal forces. This record contains present and previous values (real and imaginary) of the nodal input force loadings at each DOF.
C	OND	i	1	nnod	Ordered node list. This record exists for a load case file only.
C	OEL	i	1	nelm	Ordered element list. This record exists for a load case file only.
C	DED	i	1	sizeDead	List of dead elements (EKILL). Uses the compressed CMBLOCK format
С	MSHI	i	1	nMSHI	Mulistage harmonic indices for each stage
C C C C	ESL	i	1	2*nelm	Element solutions index table. This record contains pointers to each element solution. The order of the elements is the same as the order in the element equivalence table. (bit 12 (PDBE) in mask)
С	RCM	dp	1	6*numRotCmp	Angular velocities (3) and angular

```
C
                                        accelerations (3) of components.
    DMI
             dp
                      1 3+nContours Crack ID, Contour ID, TipNode, J Integral
C
                                        values
С
С
                                          (bit 29 (PDBCINT) in mask)
         The solution information for each individual element is stored starting
С
         at this point in the file. The next 26 records on the file are
С
         repeated as a group nelm times (once for each element). Item nelm is
C
С
         defined in the file header.
С
                                        Individual element index table.
                                        ptrEMS, ptrENF, ptrENS, ptrENG, ptrEGR,
С
                                        ptrEEL, ptrEPL, ptrECR, ptrETH, ptrEUL,
С
                                        ptrEFX, ptrELF, ptrEMN, ptrECD, ptrENL,
С
                                        ptrEHC, ptrEPT, ptrESF, ptrEDI, ptrETB,
С
                                                                                    (20)
                                        ptrECT, ptrEXY, ptrEBA, ptrESV, ptrMNL
С
С
                                        ptrESR
                                       (Relative to ptrESL)
С
                                     each item is described below:
С
                                        ptrEMS - pointer to misc. data
C
                                       ptrENF - pointer to nodal forces
C
                                        ptrENS - pointer to nodal stresses
С
                                        ptrENG - pointer to volume and energies
С
C
                                        ptrEGR - pointer to nodal gradients
                                       ptrEEL - pointer to elastic strains
ptrEPL - pointer to plastic strains
С
С
                                       ptrECR - pointer to creep strains
С
                                        ptrETH - pointer to thermal strains
C
C
                                        ptrEUL - pointer to euler angles
                                       ptrEFX - pointer to nodal fluxes
ptrELF - pointer to local forces
С
С
                                       ptrEMN - pointer to misc. non-sum values
С
С
                                        ptrECD - pointer to element current
С
                                                 densities
                                        ptrENL - pointer to nodal nonlinear data
С
С
                                        ptrEHC - pointer to calculated heat
С
                                                 generations
                                        ptrEPT - pointer to element temperatures
С
                                        ptrESF - pointer to element surface
C
С
                                                 stresses
С
                                        ptrEDI - pointer to diffusion strains
С
                                        ptrETB - pointer to ETABLE items(post1 only
                                        ptrECT - pointer to contact data
С
С
                                        ptrEXY - pointer to integration point
С
                                                 locations
                                        ptrEBA - pointer to back stresses
C
                                       ptrESV - pointer to state variables
ptrMNL - pointer to material nonlinear record
С
С
                                       ptrESR - pointer to selected results
С
                                        Note! If ptrXXX is negative, then all
C
С
                                        |ptrXXX| items are zero and are not on
                                        the file.
С
    EMS
             dр
                            varies
                                        Element summable miscellaneous data. The
С
                                        contents and number of data items is
С
                                        element-dependent. For a list of what's
C
                                        available, see the SMISC item in the
С
                                        description of the ETABLE command in the
С
                                        ANSYS Commands Reference.
C
                            varies
                                        Element nodal forces. This record contains
    ENF
                      1
С
             dρ
С
                                        the forces at each node, in the same DOF
С
                                        order as the DOF number reference table.
                                        For static, damping, and inertia forces, a
С
                                        set of forces will be repeated (as
С
С
                                        appropriate). Number of data items stored
```

c c c c c c	in this record can be calculated as follows: nodfor*NDOF*M, where NDOF is the number of DOFs/node for this element, nodfor is the number of nodes per element having nodal forces (defined in element type description record), and M may be 1, 2, or 3. For a static analysis, M=1 only. For a transient analysis, M can be 1, 2, or 3.
C ENS dp 1 varies C C C C C C C C C C C C C C C C C C C	Element nodal component stresses. This record contains the stresses at each corner node, in the order SX,SY,SY,SYX,SXZ,SXZ. Nodal order corresponds to the connectivity defined in the element description. Stresses can be nodal values extrapolated from the integration points moved to the nodes. If an element is nonlinear, integration point values always will be written. (See item rxtrap in the solution header for the setting.) An element is considered nonlinear when either plastic, creep, or swelling strains are present. Definition of common terms referred here and in subsequent EEL, EPL, ECR, ETH, ENL, EUL EPT, and EDI sections: nodstr - number of nodes per element having stresses, strains, etc. For higher-order elements, nodstr equals to the number of corner nodes (e.g., for 20-noded SOLID186, nodstr = 8). NOTE- the /config NST1 option may suppress all but one node output of nodal ENE EEL EPL ECR ETH and ENL for any given element. nodfor - number of nodes per element having nodal forces, etc. ncomp - number of solution items per node noomp = 6 for ENS record 7 for EEL record 7 for EEL record 7 for ECR record 8 for ETH record 7 for EDI record NL - number of layers in layered elements Note: For result sets with NoPrin=0, the ENS record will have ncomp=11 and include the principal stresses S1.52.53, S1,SIGE. * For solid elements or layered solid elements with KEYOPT(8)=0, the record contains stresses at each corner node and the number of items in this record is nodstr*ncomp. * For shell elements or layered shell elements with KEYOPT(8)=0, the record contains stresses at each corner node (first at the bottom shell surface, then the top surface), and the number of items in this record is 2*nodstr*ncomp. * For layered elements or layered shell elements with KEYOPT(8)=0, the record contains stresses at each corner node (first at the bottom shell surface, then the top surface), and the number of items in this record is 2*nodstr*ncomp.
c c	contains additional stresses at each corner nodes (first the bottom surface, then the
	TOURS (TITLE CITE DOCCOM BUILDING, CHEM CITE

```
top surface) of the layer with the largest
С
                                        failure criteria. Therefore, the total number
С
                                        of items is 4*nodstr*ncomp for SHELL91 and
С
С
                                        SHELL99, and 2*nodstr*ncomp for SOLID46 and
С
                                        SOLID191.
                                       * For layered elements (with KEYOPT(8)=1),
C
                                        stresses for each layer are at each
С
                                        corner node (first at the bottom surface, then
С
                                        at the top surface), and the number of
C
С
                                        items in this record is NL*2*nodstr*ncomp for
                                        layered shells and NL*nodstr*ncomp for
C
                                        layered solid elements.
С
                                       * For layered shell elements with KEYOPT(8)=2,
С
                                        the record contains stresses for each layer
C
                                        at each corner node (first at the bottom
С
                                        surface, then the top, and finally the middle
С
                                        surface). Therefore, the number of items
С
                                        in this record is NL*3*nodstr*ncomp.
С
                                      * For layered membrane elements (SHELL181,
С
                                        SHELL281, SHELL208, and SHELL209 with
C
                                        KEYOPT(1)=1 and KEYOPT(8)=1), the record
C
                                        contains stresses for each layer at each
C
С
                                        corner node, and the number of items in
                                        this record is NL*nodstr*ncomp.
С
                                       * For beam elements, the contents and number
C
                                        of data items is element-dependent. See
С
С
                                        the Output Data section for the particular
C
                                        element in the ANSYS Elements Reference.
С
    ENG
                           varies
                                      Element volume and energies.
            dρ
C
                                      * if cpxeng = 0, lrec = 11:
C
С
                                        volu, sene, aene, kene, coene,
С
                                      incene.
                                                0.0,
                                                        0.0, thene,
                                                                         0.0, (10)
C
С
                                         0.0
                                      * if cpxeng = 1, lrec = 21:
C
С
                                                                kene, coene,
С
                                        volu, sene,
                                                        aene,
                                      incene, 0.0,
                                                        0.0, thene,
                                                                         0.0, (10)
C
                                         0.0, dene, wext, asene, akene,
C
С
                                       psene, pkene, asint1, akint1, asint2, (20)
С
                                      akint2
                                    each item is described below:
C
С
                                      volu
                                             - element volume
                                             - element energy associated with
                                      sene
C
                                                the stiffness matrix
С
                                             - artificial hourglass energy
С
                                      aene
                                            - kinetic energy
С
                                      kene
                                      coene - co-energy (magnetics)
C
                                      incene - incremental energy (magnetics)
C
                                             position not usedposition not used
С
                                      0.0
С
                                      0.0
                                      thene - thermal dissipation energy
C
С
                                                (see ThermMat, shell131/132 only)
                                      0.0
                                             - position not used
С
                                             - position not used
                                      0.0
C
                                             - damping energy
С
                                      dene
                                             - work due to external element load
С
                                      wext.
                                      asene - amplitude stiffness energy
C
С
                                      akene - amplitude kinetic energy
                                      psene - peak stiffness energy
C
                                      pkene - peak kinetic energy
asint1 - first intermediate result
С
С
                                                for asene calculations
C
                                      akint1 - first intermediate result
С
С
                                                for akene calculations
```

с с с				<pre>asint2 - second intermediate result for asene calculations akint2 - second intermediate result for akene calculations</pre>
C EGR C C C C C C C C C C C C C C C C C C C	dp	1	varies	Element nodal field gradients. This record contains the gradients at each corner node in the order X,Y,Z. Nodal order corresponds to the connectivity defined in the element description. If this is a coupled-field analysis, the data is stored in the following order (as available): fluid, thermal (TEMP), electric (VOLT), magnetic (AZ), and diffusion (CONC). Gradients can be nodal values extrapolated from the integration points or values at the integration points moved to the nodes. See item rxtrap in the solution header for the setting. The number of items in this record is nodstr*3*N, where N can be 1, 2, 3, or 4 (depending on the coupled-field conditions). NOTE: nodstr is defined in the element type description record.
C EEL C C C C C C C C C C C C C C C C C C	dp	1	varies	Element nodal component elastic strains. This record contains strains in the order X,Y,Z,XY,YZ,XZ,EQV. Elastic strains can be can be nodal values extrapolated from the integration points or values at the integration points moved to the nodes. If an element is nonlinear, integration point values always will be written. See item rxtrap in the solution header for the setting. An element is considered nonlinear when either plastic, creep, or swelling strains are present. For beam elements, see item LEPEL in the description in the Output Data section for the particular element in the ANSYS Elements Reference. NOTE: See ENS record section for more details on record content and length.
C EPL C C C C C C C C C	dp	1	varies	Element nodal component plastic strains. This record contains strains in the order X,Y,Z,XY,YZ,XZ,EQV. Plastic strains are always values at the integration points moved to the nodes. For beam elements, see item LEPPL in the Output Data section for the particular element in the ANSYS Elements Reference. NOTE: See ENS record section for more details on record content and length.
C ECR C C C C C C C C	dр	1	varies	Element nodal component creep strains. This record contains strains in the order X,Y,Z,XY,YZ,XZ,EQV. Creep strains are always values at the integration points moved to the nodes. For beam elements, see item LEPCR in the Output Data section for the particular element in the ANSYS Elements Reference. NOTE: See ENS record section for more details on record content and length.

```
C
    ETH
            dp
                     1
                           varies
                                      Element nodal component thermal strains.
C
                                      This record contains strains in the order
С
С
                                      X,Y,Z,XY,YZ,XZ,EQV plus the element
                                      swelling strain. Thermal
C
                                      strains can be nodal values extrapolated
С
                                      from the integration points or values at
С
                                      the integration points moved to the nodes.
C
C
                                      If the element in nonlinear, integration
                                      point data always will be written. (An
C
                                      element is considered nonlinear when either
С
C
                                      plastic, creep, or swelling strains are
                                      present.) See item rxtrap in the solution
C
                                      header for the setting. For beam elements,
С
С
                                      see item LEPTH in the description of the
                                      Output Data section for the particular
С
                                      element in the ANSYS Elements Reference.
С
С
                                      NOTE: See ENS record section for more details
С
С
                                      on record content and length.
C
    EUL
            dр
                     1
                          varies
                                      Element Euler angles. This record contains
С
                                      the Euler angles (THXY, THYZ, THZX). (No
C
                                      attempt is made to make this information
С
С
                                      complete for all cases of all element
C
                                      types. Programmers need to verify their
С
                                      situations.)
С
                                      ** FOR OLDER ELEMENTS **
С
                                      --For lower-order elements
C
                                        (e.g. PLANE42 and SOLID45), angles are
C
                                        at the centroid and the number of items
C
С
                                        in this record is 3.
                                      --For higher-order elements (e.g.
С
С
                                        PLANE82, SOLID92, and SOLID95), angles
С
                                        are at each corner node and the number of
                                        items in this record is nodstr*3.
C
С
                                        The above two categories are output if
                                        ESYS is used, a material KEYOPT is used
С
                                        (e.g. KEYOPT(1) for PLANE42), or if it is
С
                                        from a large deflection superelement
C
С
                                        stress pass
С
С
                                      ** FOR NEW GENERATION SHELL ELEMENTS **
                                      --For SHELL181/SHELL281:
С
С
                                         For real constant input: the number of
С
                                           items in this record is 12.
                                         For section input:
C
                                          IF KEYOPT(8) > 0:
С
                                           the number of items in this record is
С
С
                                                            12 + number of layers
                                          IF KEYOPT(8) = 0:
C
                                             IF regular shell (KEYOPT(1) = 0)
C
                                                the number of items in this record
С
С
                                                 is 14
                                            IF membrane shell (KEYOPT(1) = 1):
C
                                               IF number of layers = 1, the number
С
                                                of items in this record is 13
С
                                               IF number of layers > 1, the number
C
С
                                                of items in this record is 14
С
                                      ** FOR THE NEW GENERATION SOLID ELEMENTS **
C
С
                                      --For uniform reduced integration lower-order
                                        elements (e.g. PLANE182, KEYOPT(1)=1 and
C
С
                                        SOLID185 KEYOPT(2)=1):
С
                                        the angles are at the centroid and the number
                                        of items is 3.
C
                                       --For other formulations of lower-order
С
С
                                        elements (e.g. PLANE182 and SOLID185) and
```

C					the higher-order elements
С					(e.g. PLANE183, SOLID186, and SOLID187):
С					The number of items in this record is
C					<pre>(nodstr*3)For layered solid elements, add NL values,</pre>
C					so that the number of items in this record
C					is (nodstr*3)+NL.
C					IS (HOUSEL S) THE.
С					NOTE: See ENS record section for definition of
C					terms NL and nodstr.
С	EFX	dp	1	varies	Element nodal field fluxes. This record
С					contains the fluxes at each corner node in
C					the order X,Y,Z. If this is a
C					coupled-field analysis, the flux data is
С					stored in the following order: thermal,
С					electric, magnetic. Nodal order
С					corresponds to the connectivity defined in
C					the element description. Fluxes can be nodal values extrapolated from the
C					integration points or values at the
C					integration points moved to the nodes.
С					See item rxtrap in the solution header for
C					the setting. The number of items in this
С					record is nodstr*3*N, where N can be 1, 2,
С					or 3 depending on the coupled-field
C					conditions.
С					NOTE: nodstr is defined in the element type
C					description record.
			1		
	ELF	dp	Τ	varies	Element nodal coupled-field forces. This
С					record lists the forces at each node in the
C					order X,Y,Z. For most elements, the number of items in this record is nodfor*3.
C					However, for the PLANE53 element, the
C					number of items in this record is either
C					nodfor*3 or nodstr*3. (See the description
С					of KEYOPT(7) for PLANE53 in the ANSYS
С					Elements Reference.) NOTE: nodfor and
С					nodstr are defined in the element type
С					description record.
C	EMN	dp	1	varies	Element nonsummable miscellaneous data.
С					The contents and number data items for this
C					record is element-dependent. See the
С					description for item NMISC of the ETABLE
С					command in the ANSYS Commands Reference.
~ +	ECD	dn	1	3	Element current densities. This record
c ^	FCD	dp	1	3	contains the calculated current densities
C					in the order X,Y,Z.
С	ENL	dp	1	varies	Element nodal nonlinear data. This record
С		_			stores nonlinear data at each corner node
C					in the order SEPL, SRAT, HPRES, EPEQ,
С					PSV or CREQ, PLWK, CRWK, and ELENG
C					followed by 2 spares.
С					each item is described below:
С					SEPL - equivalent stress parameter
C					SRAT - stress ratio
С					HPRES - hydrostatic pressure
C					EPEQ - accumulated equivalent plastic strain
C					PSV - plastic state variable
C					CREQ - accumulated equivalent creep
С					strain. Applies to current
С					technology element types
C					180,181,182,183,185,186,
C					187,188,189,208,209,265,

C					281,288,289,290
С					PLWK - plastic strain energy density(work)
С					CRWK - creep strain energy density (work)
С					ELENG - elestic strain energy density
С					* See ENS record section for details on
С					solid and shell elements.
С					* For beam elements, the contents and
C					number of data items in this record is
С					element-dependent. See the description
C					of item NLIN in the Output Data section
С					for the particular element in the ANSYS
					Elements Reference.
С					Elements Reference.
_ +	EHIO	a	1	1	Dlawart hast consention White warrand
	EHC	dp	1	1	Element heat generation. This record
С					stores the calculated heat generation.
_	n.p.m	a	1		71
С	EPT	dp	1	varies	Element structural nodal temperatures.
С					* For solid elements and SHELL41, the
С					record contains nodal temperatures at
С					each node and the number of items in this
C					record is nodfor.
C					* For shell elements, except SHELL41 and
C					SHELL91, the record contains nodal
C					temperatures at each corner node for the
C					top surface and the bottom surface. The
C					number of items in this record is
C					nodstr*2.
C					* For SHELL91 and SOLID191, the record
C					contains nodal temperatures at each
C					corner node for the bottom of the bottom
С					layer, and each succeeding interlayer
С					surface up to the top of the top layer.
С					The number of items in this record is
С					(NL+1)*nodstr.
С					* For layered shell elements SHELL181,
С					SHELL281, SHELL208, SHELL209, and layered
С					solid elements SOLID185, SOLID186,
С					and SOLSH190, the record contains
С					temperatures for each layer at each
C					corner node (first at the bottom layer
С					surface, then the top). Therefore, the number
C					of items in this record is NL*2*nodstr for
С					layered shells and NL*nodstr for layered
C					solid elements. * For laward membrane elements (SHFII181
C					* For layered membrane elements (SHELL181,
C					SHELL281, SHELL208, and SHELL209 with
C					KEYOPT(1)=1), the record contains
С					temperatures for each layer at each
C					corner node. Therefore, the number of items
С					in this record is NL*nodstr.
С					* For beam elements, the contents and
С					number of data items in this record is
C					element-dependent. See the description
C					of item LBFE in the Output Data section
C					for the particular element in the ANSYS
C					Elements Reference.
С					NOTE: See ENS record section for definition
C					of terms NL, nodstr, and nodfor.
C	ECT	dp	1	varies	Contact element results. This record
C					stores contact results at each corner node
C					in the order STAT, PENE, PRES, SFRIC, STOT,
C					SLIDE, GAP, FLUX, CNOS, FPRS
C					each item is described below:
C					STAT - Contact status
С					PENE - Contact penetration
С					PRES - Contact pressure
C					SFRIC - Contact friction stress

```
STOT - Contact total stress (pressure plus friction)
С
                                     SLIDE - Contact sliding distance
С
                                     GAP - Contact gap distance
С
                                     FLUX - Total heat flux at contact surface
С
С
                                     CNOS - Total number of contact status changes during substep
                                     FPRS - Actual applied fluid penetration pressure
C
    ESF
                         nsurf*19
                                     Element surface stresses. The
С
            dр
                     1
                                     length of this record is nsurf*19 where
C
С
                                     nsurf is the number of surfaces that have
                                     surface stress information. The stress
C
                                     information is simply repeated in the
С
С
                                     format shown below for each surface.
                                     * For 2d elements:
С
С
                                     facenm, area, temp, press, eppar,
                                              epz, 0.0d0, spar, sper,
С
                                      epper,
С
                                         sz, 0.0d0, 0.0d0, 0.0d0,
                                               s3, sint, seqv
C
                                     * For 3d elements:
C
                                     facenm, area, temp, press,
С
                                              epz, epxy, sx,
C
                                        еру,
                                                                     sy,
                                               sxy, 0.0d0, 0.0d0,
С
                                         SZ,
                                                                     s1,
С
                                         s2,
                                              s3, sint, seqv
                                     * For axisymmetric elements:
С
                                     facenm, area, temp, press, eppar,
C
                                      epper, epz, epsh, spar, sper,
C
                                         sz, 0.0d0, 0.0d0, ssh,
C
                                                                   s1,
                                         s2,
                                               s3, sint, seqv
С
                                   each item is described below:
C
С
                                     facenm - face number
                                     area - face area
C
                                            - face temperature
С
                                     temp
                                     press - face pressure
С
                                            - strain parallel to face
С
                                     epx
                                            - strain parallel to face
С
                                     еру
С
                                     epz
                                            - strain perpendicular to face
                                            - shear strain
С
                                     ерху
                                     eppar - strain parallel to face
С
                                     epper - strain perpendicular to face
С
С
                                     epsh
                                           - torsion shear strain
С
                                     SX
                                            - stress parallel to face
                                            - stress parallel to face
C
                                     sy
                                            - stress perpendicular to face
С
                                     SZ
                                            - shear stress
С
                                     SXY
                                           - stress parallel to face
С
                                     spar
                                           - stress perpendicular to face
С
                                     sper
                                     ssh
                                            - torsion shear stress
C
С
                                     s1
                                            - S(1)
                                            - S(2)
С
                                     s2
                                            - S(3)
                                     s3
C
                                     sint - S(INT)
С
С
                                     seqv
                                            - S(EQV)
                                     0.0d0 - position not used
C
    EDI
                    1
                                     Element nodal component diffusion strains.
С
           dр
                          varies
                                     This record contains strains in the order
C
С
                                     X,Y,Z,XY,YZ,XZ,EQV. Diffusion
                                     strains can be nodal values extrapolated
С
С
                                     from the integration points or values at
С
                                     the integration points moved to the nodes.
                                     See item rxtrap in the solution header for
C
                                     the setting.
С
```

c NOTE: See ENS record section for more details	
c on record content and length.	
C EXY dp 1 varies Element integration point coordinates The length of the record is numint*3, where numint is the number of integration points. Even two-dimensional elements use the 3. They are output only if requested with the OUTRES,loci command. Applicable only to legacy element types 2,42,45,82,92,95, and current technology element types 180,181,182,183,185,186,187, 188,189,208,209,265,281,288,289,290	
C EBA dp 1 varies Element structural nodal back stresses Record has the same form as the plastic strains. They are output if the form of plasticity is kinematic hardening and the plastic strains are requested. Applicable only to legacy element types 2,42,45,82,92,95, and current technology element types 180,181,182,183,185,186,187, 188,189,208,209,265,281,288,289,290	
c ESV dp 1 varies Element state variable record. Exists only c if written by user in usermat or usercreep.	
c MNL dp 1 varies Material nonlinear record.	
c ESR dp 1 varies Element nodal selected results. The length	
c is NINT(ESR(1)). The record includes c type specifications and results values. c The type of results depends on the selected c result output specfications.	
c type specifications and results values. c The type of results depends on the selected	
c type specifications and results values. c The type of results depends on the selected result output specifications. c records marked with * to the left of the record id can be read and stored	
type specifications and results values. The type of results depends on the selected result output specifications. records marked with * to the left of the record id can be read and stored into database with "ldread" command. The solution information for the Nodal Averaged Results (NAR) is stored	
type specifications and results values. The type of results depends on the selected result output specifications. records marked with * to the left of the record id can be read and stored into database with "ldread" command. The solution information for the Nodal Averaged Results (NAR) is stored starting at this point in the file.	
type specifications and results values. The type of results depends on the selected result output specifications. c records marked with * to the left of the record id can be read and stored into database with "ldread" command. c The solution information for the Nodal Averaged Results (NAR) is stored starting at this point in the file. c NAR i 1 10 NAR index table. c ptrNCTl, ptrNCTh, ptrNSTl, ptrNSTh, ptrNELl, ptrNELh, ptrNPLl, ptrNPLh, ptrNPLh, ptrNPLh, ptrNCRh, (10) ptrNTHl, ptrNTHh	
type specifications and results values. The type of results depends on the selected result output specifications. c records marked with * to the left of the record id can be read and stored into database with "ldread" command. c The solution information for the Nodal Averaged Results (NAR) is stored starting at this point in the file. c NAR i 1 1 0 NAR index table. c ptrNCTl, ptrNCTh, ptrNSTl, ptrNSTh, ptrNELl, ptrNELh, ptrNELh, ptrNELh, ptrNPLh, ptrNPLh, ptrNCRl, ptrNCRh, (10) ptrNTHl, ptrNTHh c (Relative to ptrNAR) c NOTE: nodal results are stored in the same node order as given above in the nodal equivalence table. If a DOF for a node isn't valid, a value of 2.0**100	
type specifications and results values. The type of results depends on the selected result output specifications. crecords marked with * to the left of the record id can be read and stored into database with "ldread" command. cThe solution information for the Nodal Averaged Results (NAR) is stored starting at this point in the file. cNAR i 1 10 NAR index table. cPrinctl, ptrnCth, ptrnStl, ptrnSth, ptrnEtl, ptrnEth, ptrnPLL, ptrnPLL, ptrnPLL, ptrnPLL, ptrnThL, ptrnThL, ptrnThL, ptrnThL, ptrnThL, ptrnThL, ptrnThL, compared to the condition of the same node order as given above in the nodal equivalence table. If a DOF for a node isn't valid, a value of 2.0**100 is used. cNCT i 1 4*nnod Node Contributions. This record contains the number of elements that shared each node and contributed to the calculated value for each NAR vector. The first nnod values are the node contributions for he nodal stresses, the next nnod values are for the node contibutions for nodal elastic	

```
C
                                      node, in the order X,Y,Z,XY,YZ,XZ,EQV.
    NPL
            dp
                           7*nnod
                                      Nodal component plastic strains.
C
                                      This record contains the strains at each
С
С
                                      node, in the order X,Y,Z,XY,YZ,XZ,EQV.
    NCR
                           7*nnod
                                      Nodal component creep strains.
С
            dр
                                      This record contains the strains at each
С
                                      node, in the order X,Y,Z,XY,YZ,XZ,EOV.
C
    NTH
                     1
                           8*nnod
                                      Nodal component thermal and swelling
С
            dρ
С
                                      strains. This record contains the strains
С
                                      at each node, in the order X,Y,Z,XY,YZ,XZ,
                                      EQV plus the node swelling strain
C
c *** Nodal Component Stresses (unused)
С
  NDSTR
                     1
                           6*nnod
                                      Nodal component stresses (TOP for shells)
                                      (nNodStr > 0)
C
                           6*nnod
                                      BOT nodal component stresses for shells
С
                                      (nNodStr > 1)
C
С
                          6*nnod
                                      MID nodal component stresses for shells
                                       (nNodStr > 2)
```

1.3. Description of the Reduced Displacement File

This section explains the content of the reduced displacement file (jobname.rdsp).

1.3.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 10.

1.3.2. RDSP File Format

```
*comdeck,fdrdsp
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc
     ******* description of reduced displacement file *******
     character*8 RDSPNM
     parameter (RDSPNM='rdsp
     integer
                    RDSPHDLEN
     parameter
                   (RDSPHDLEN=80)
     LONGINT
                    rdspfpL, rdspfp
                    rdspbk, rdsput
     common /fdrdsp/ rdspfpL, rdspbk, rdsput
     equivalence (rdspfp,rdspfpL)
 write: lnfrcl,lnfrin,lnfrwr
 read:
         rdtrs
   ****** common variable descriptions ********
C
  rdspfpL
               file position on file rdsp
CO
   rdspbk
                block number for file rdsp
               file unit for file rdsp
CO
   rdsput
```

```
C
    See fddesc for documentation of how binary files are stored.
С
      ****** file format ******
C
С
         recid tells the identifier for this record. Not all records will have
              identifiers -- they are only indicated for those records whose
С
              record pointers are stored in the second file header.
С
         type tells what kind of information is stored in this record:
C
С
              i - integer
              dp - double precision
C
              cmp - complex
С
         nrec tells how many records of this description are found here
C
         lrec tells how long the records are (how many items are stored)
c recid
           type
                   nrec
                           lrec
                                    contents
                    1
                           100
                                    standard ANSYS file header (see binhed8 for
С
                                    details of header contents)
C
                    1
                            80
                                    .RDSP FILE HEADER
С
С
                                 fun10,
                                                              nmode, numdof,
C
                                           nmrow, nmatrx,
С
                                  maxn,
                                           wfmax, lenbac, ngaps, ncumit, (10)
                                                   ndva,
                                                             nvect, DSPfmt,
С
                                   kan,
                                          nres,
C
                                minmod.
                                             0, modlstp, ndefdval, nEnfDof, (20)
                                                                   0,
                                ptrDOF, ptrDAMP, ptrDAMPh,
С
                                                                              0,
                                                        0,
С
                                ptrFRQ,
                                         ptrDSP,
                                                                    Ο,
                                                                              0, (30)
                               ptrFRQh, ptrDSPh,
                                                    ptrDVA, ptrDVAh,nrkeyPert,
C
                              kPerturb,
                                           keyVA,Glblenbac,
                                                                   0,
                                                                              0, (40)
C
                                               0,
                                                        0,
                                                                   0,
                                                                             0,
                                     0,
C
                                     0,
                                               0,
                                                         0,
                                                                    0,
                                                                              0, (50)
С
                                                        0,
С
                                     0,
                                               0,
                                                                   0,
                                                                             0,
                                                        0,
                                                                   0,
                                     0,
                                               0,
                                                                             0, (60)
C
                                                         0,
С
                                     0,
                                               0,
                                                                    0,
                                                                             Ο,
С
                                     Ο,
                                               0,
                                                         0,
                                                                    Ο,
                                                                             0, (70)
                                                         0,
                                     0.
                                               0,
                                                                   0,
                                                                             0,
С
С
                                     0,
                                               0,
                                                         0,
                                                                    0,
                                                                              0 (80)
                                 each item in header is described below:
С
С
                                   fun10 - unit number (rdsp file is 10)
                                   nmrow - number of rows/columns in matrices
С
С
                                   nmatrx - number of reduced matrices on the
                                            file
C
С
                                   nmode - number of modes extracted during
С
                                           modal analysis
                                   numdof - number of dofs per node
C
                                   maxn - maximum node number
С
                                   wfmax - maximum wavefront
С
                                   lenbac - number of nodes
С
                                   ngaps - number of gaps
C
                                   ncumit - total number of iterations done
C
С
                                            during analysis
                                          - analysis type
С
                                   kan
                                            = 5 for MSUP transient analysis
C
                                   nres - number of residual vectors used
С
С
                                   modlstp- multiple load step key
                                   ndva - length of DVA (for restart)
C
С
                                   nvect - number of available load vectors
                                   DSPfmt - 0,physical disps .ne.0,modal coords
С
                                   minmod - smallest mode number used
C
С
                                   ndefdval- number of defined enforced motion
                                   maxEnf - maximum enforced values
С
                                   ptrDOF - pointer to degree of freedom set
ptrDAMP - pointer to damping values
С
С
                                   ptrDAMPh - High part of DAMP pointer
C
                                   ptrDNC - pointer to nodal constraints
С
С
                                   ptrFRQ - pointer to the frequencies
```

```
С
                                    ptrDSP - pointer to the step solution data:
С
                                                - calculated displacements
                                                - load vector scale factors
C
С
                                                - gap restoring forces
С
                                                - calculated velocities
                                                - calculated accelerations
C
                                    ptrFRQh- High part of frequency ptr
С
                                    ptrDSPh- High part of displacement ptr
С
                                    ptrDVA - pointer to modal disp, velo and acc (for restart)
C
C
                                    ptrDVAh- High part of modal disp, velo and acc (for restart)
                                    nrkeyPert - nrkey setting of base analysis (Linear Perturbation)
С
С
                                    kPerturb - Linear Perturbation key
С
                                    keyVA - Key for velocities/accelerations on file
                                    Glblenbac - global number of nodes (== 0 unless using Distributed Ansys)
C
С
                                           - position not used
С
            i
                    1
                          numdof
                                    Degrees of freedom per node
                                     (curdof(i),i=1,numdof)
С
С
                                    dof reference numbers are:
          UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8
C
             = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
C
          \texttt{CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG=22, ENKE=23, ENDS=24}
С
С
          EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
                           lenbac Nodal equivalence table. This table equates
C
С
                                     the number used for storage to the actual
С
                                     node number
                                    (Back(i), i=1, lenbac)
C
                    1 Glblenbac Global nodal equivalence table. This
C
                                     table equates the number used for storage
C
                                     to the actual node number. Only written
C
C
                                     by the master process in Distributed Ansys
                                    (GlbBack(i), i=1,Glblenbac)
С
                            10
                                   Time information:
            ďρ
                    1
C
С
                                    dtime,
                                              0.0,
                                                       0.0,
                                                               0.0,
                                                                       0.0,
С
                                      0.0,
                                              0.0,
                                                      0.0,
                                                               0.0, timend
                                  each item is described below:
С
С
                                    dtime - the time increment
С
                                    timend - the final time of the analysis
C
С
                                         - position not used
С
   DOF
            i
                    1
                          nmrow
                                    Degree of freedom set used
                                    The DOFs are calculated as (N-1)*numdof+DOF,
C
С
                                    where N is the position number of the node in
С
                                    the nodal equivalence table and DOF is the
                                    DOF reference number given above.
C
                                   Original reduced set of DOFs used.
С
                    1
                          nmrow+1
                                    The DOFs are calculated as (N-1)*numdof+DOF,
C
                                    where N is the position number of the node in
C
                                    the nodal equivalence table and DOF is the
C
С
                                    DOF reference number given above.
C
   DAMP
            dp
                        nmode+10
                                    Damping values.
С
                                    This record is present only for analysis using
С
                                    the mode superposition method with MCkey
                                    activated - TRNOPT cmd.
C
С
                                    There are nmode+10 entries:
                                                     - effective damping ratios
С
                                    1 : nmode
                                    nmode+1: nmode+5 - alphad, betad, dmprat, dmpstr, dsmpFreq
C
                                    nmode+6 : nmode+10 - additional entries - zeroed out for now
С
   FRQ
            dр
                    1
                           nmrow/
                                    Frequencies extracted from the modal analysis.
C
С
                           ndva/
                                    This record is present only for analyses using
                      ndva-nEnfDof the mode superposition method.
С
                                    If DSPfmt is 0, the first nmrow values are the
С
С
                                    frequencies extracted from the modal analysis.
```

```
C
                                    The remaining values have no meaning. Otherwise,
С
                                    there are ndva frequencies if there are no
                                    enforced motions and ndva less the number of enforced
C
С
                                    motions otherwise.
С
                                    If DSPfmt=0 :(freq(i),i=1,nmrow)
                                    If DSPfmt!=0 and ndefdval=0:
C
                                      (freq(i),i=1,ndva)
С
                                    If DSPfmt!=0 and ndefdval>0:
С
                                      (freq(i),i=1,ndva-nEnfDof)
C
   *** The next 6 to 7 records are repeated (as a group) until the time value
C
   *** equals the value of timend. The number of iterations is stored as
С
С
   *** ncumit. (see above records that deal with time)
                           nmrow+7/ Calculated displacements
С
   DSP
            dр
С
                          ndva+7
                                   The first nmrow entries are the displacements
                                    in the same order as the original set of DOFs
C
                                    (see record AFTER ptrDOF). If DSPfmt=0, these
С
С
                                    are physical displacements, If DSPfmt!=0,
                                    these are the ndva modal coordinates instead
С
                                    of the nmrow entries.
C
C
С
                                    For the last six entries:
                                     1. Time for these displacements
С
                                     2. Load step number
C
                                     3. Substep number
С
                                     4. Cumulative iteration number
С
С
                                     5. Scale factor
                                     6. numdeflys - number of scale factors
С
                                     7. kwrval - key to control writing velocities
С
                                               and accelerations at each substep (write if keyVA = 1 and kwrval =
С
С
С
            i
                    1
                                    Note: If, upon reading of this record, there
                                    is less than nmrow+5 items in the record,
С
С
                                    then only a selected set of nodes were
                                    output. Another record follows (integer, less
С
С
                                    than lenbac long) which contains the list of
С
                                    nodes for which DOF solutions are available.
С
            i
                    1 numdeflvs
                                    lvscal table scale factor IDs
                                     (ilvscID(i), i=1, numdeflvs)
С
                    1 numdeflvs
                                    lvscal table scale factor values
C
            dp
С
                                     (dlvscVal(i),i=1,numdeflvs)
                                    Gap restoring forces. This record is
C
            dp
                           ngaps
                                    present only if ngaps > 0.
C
С
                                     (fgaps(i),i=1,ngaps)
                                    Calculated velocities (present if keyVA = 1).
                    1 nmrow/ndva
C
            dp
                                    nmrow entries if DSPfmt=0, ndva entries if DSPfmt!=0
С
            dp
                    1 nmrow/ndva
                                    Calculated accelerations (present if keyVA = 1)
C
                                    nmrow entries if DSPfmt=0, ndva entries if DSPfmt!=0
c *** The next 3 records are kept for possible restart in mode superposition
c *** transient. They are overwritten upon restarting. They are written once (last
c *** loadstep).
С
  DVA
            ďρ
                    1
                          ndva+6 Calculated modal displacements
                                    The first ndva entries are the modal
C
С
                                    displacements. For the last six entries:
                                     1. Time for these displacements
С
                                     2. Load step number
C
С
                                     3. Substep number
                                     4. Cumulative iteration number
С
С
                                     5. Scale factor
C
                                     6. numdeflys - number of scale factors
                 1 ndva Calculated modal velocities
```

```
c --- dp 1 ndva Calculated modal accelerations
```

1.4. Description of the Reduced Complex Displacement File

This section explains the content of the reduced complex displacement file (jobname.rfrq).

1.4.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 10.

1.4.2. RFRQ File Format

```
*comdeck,fdrfrq
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc
      ****** description of reduced complex displacement file *******
      character*8 RFRQNM
      parameter (RFRQNM='rfrq
     LONGINT
                    rfrqfpL, rfrqfp
                    rfrqbk, rfrqut
      common /fdrfrq/ rfrqfpL, rfrqbk, rfrqut
      equivalence (rfrqfp,rfrqfpL)
   ****** common variable descriptions ********
C
co rfrqfpL file position on file rfrq
co rfrqbk
                block number for file rfrq
co rfrqut
                file unit for file rfrq
   See fddesc for documentation of how binary files are stored.
C
С
      ****** file format *******
С
С
         recid tells the identifier for this record. Not all records will have
             identifiers -- they are only indicated for those records whose
C
             record pointers are stored in the second file header.
C
         type tells what kind of information is stored in this record:
C
С
             i - integer
С
             dp - double precision
             cmp - complex
C
        nrec tells how many records of this description are found here
         lrec tells how long the records are (how many items are stored)
c recid
                          lrec
                                   contents
          type
                 nrec
                   1
                          100
                                   standard ANSYS file header (see binhed8 for
C
                                    details of header contents)
С
C
                   1
                           40
                                  .RFRO FILE HEADER
С
                                     fun10.
                                              nmrow, nmatrx, nmode, numdof,
С
                                             wfmax, lenbac, extopt, ncumit, (10)
nres, nmUsed, nvect, DSPfmt,
                                     maxn,
C
                                            nres, nmUsed, nvect, DSPfmt,
0, modlstp, 0, nEnfdof, (20)
C
                                    minmod,
C
                                    ptrDOF, ptrDAMP, ptrDAMPh,
                                                                      0,
```

```
С
                                     ptrFRQ, ptrDSP,
                                                               0,
                                                                         0,
                                                                                   0, (30)
                                    ptrFRQh, ptrDSPh,nrkeyPert, kPertrb,Glblenbac,
С
                                     cpxmod.
                                               SvCode, QRdampKey,
                                                                         0,
                                                                                    0 (40)
C
С
                                  each item in header is described below:
                                    fun10 - unit number (rfrq file is 10)
С
                                    nmrow - number of rows/columns in matrices
С
                                    nmatrx - number of reduced matrices on file
C
C
                                    nmode - number of modes extracted during
                                             modal analysis
C
                                    numdof - number of dofs per node
С
C
                                    maxn - maximum node number
                                    wfmax - maximum wavefront
C
                                    lenbac - number of nodes
С
                                    extopt - mode extraction method
С
                                              = 3 - unsymmetric Lanczos
С
                                              = 4 - damped Lanczos
С
                                              = 6 - block Lanczos
С
                                              = 8 - SuperNode
С
                                              = 9 - PCG Lanczos
C
                                    ncumit - total number of iterations done
C
С
                                             during analysis
                                            - analysis type
С
                                             = 6 - MSUP harmonic analysis
C
                                    nmUsed - number of modes used in mode
С
С
                                             superposition
                                    nvect - number of generated loads in .mlv
C
                                    DSPfmt - 0,physical disps .ne.0,modal coords
С
                                    minmod - smallest mode number used
С
                                    modlstp- multiple load step key
С
                                    ptrDOF - pointer to degree of freedom set
C
C
                                             used in model
                                    ptrDAMP - pointer to damping values ptrDAMPh - High part of DAMP pointer
С
С
                                    ptrFRQ - pointer to the frequencies
С
С
                                    ptrDSP - pointer to the calculated
С
                                             displacements
                                    ptrFRQh- High part of FRQ pointer
С
С
                                    ptrDSPh- High part of DSP pointer
                                    nrkeyPert - nrkey setting of base analysis (Linear Perturbation)
С
                                    kPertrb- Linear Perturbation key
C
                                    Glblenbac - global number of nodes (== 0 unless using Distributed Ansys)
C
C
                                    cpxmod - key if complex modes were used
С
                                    SvCode - Solver assembly code
С
                                             = 1 Symbolic assembly
                                    QRdampKey - Key for QRDAMP solver use
С
С
                                                for modal analysis preceeding MSUP (MODOPT, QRDAMP)
С
                                               = 1 if QRDAMP was used, 0 otherwise
C
                                           - position not used
С
            i
                    1
                          numdof
                                   Degrees of freedom per node
C
                                     (curdof(i),i=1,numdof)
С
                                    dof reference numbers are:
C
          UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8
С
          AZ = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
C
          CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
C
          EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
                           lenbac Nodal equivalence table. This table equates
C
С
                                     the number used for storage to the actual
                                     node number
С
                                    (Back(i), i=1, lenbac)
C
                    1 Glblenbac Global nodal equivalence table. This
            i
C
С
                                     table equates the number used for storage
                                     to the actual node number. Only written
С
                                     by the master process in Distributed Ansys
C
                                    (GlbBack(i), i=1,Glblenbac)
C
```

```
C
            ďρ
                             10
                                   Unused record. contents:
С
                                   1.0, 0.0, 0.0, 0.0, 0.0,
                                   0.0, 0.0, 0.0, 0.0, 0.0
C
С
  DOF
            i
                    1
                           nmrow
                                    Degree of freedom set used
                                    The DOFs are calculated as (N-1)*numdof+DOF,
C
                                    where N is the position number of the node in
С
                                    the nodal equivalence table and DOF is the
С
                                    DOF reference number given above.
C
                           nmrow+1
                                    Original reduced set of DOFs used.
            i
                    1
C
С
                                    The DOFs are calculated as (N-1)*numdof+DOF,
С
                                    where N is the position number of the node in
                                    the nodal equivalence table and DOF is the
C
                                    DOF reference number given above.
С
С
   DAMP
            dp
                    1
                        nmode+10
                                    Damping values.
                                    This record is present only for analyses using
С
С
                                    the mode superposition method with MCkey
                                    activated - HROPT cmd.
С
                                    There are nmode+10 entries:
C
                                    1 : nmode - effective damping ratios
C
С
                                    nmode+1 : nmode+4 - alphad, alphad, dmprat, dmpstr
                                    nmode+5 : nmode+10 - additional entries - zeroed out for now
С
                                    Frequencies extracted from the modal analysis.
С
   FRO
            dр
                           nmrow/
С
                        nmode+nres
                                    This record is present only for analyses using
                                    the mode superposition method.
С
С
                                    There are nmrrow entries if DSPfmt=0 and nmode+nres
                                    entries otherwise.
С
                                    If cpxmod
С
                                    If DSPfmt=0 : (freq(i),i=1,nmrow)
C
                                    If DSPfmt!=0 : (freq(i),i=1,nmode+nres)
C
                                    Frequencies are complex if DSPfmt=0 and extopt=3 or 4
С
С
                                    or if DSPfmt!=0 and cpxmod = 1.
                                    For complex frequencies the number of entries is doubled.
C
С
                                    Real parts appear for first half and imaginary parts next.
С
                                    If DSPfmt!=0 and cpxmod = 1:
                                    (dreal(freq(i)),i=1,nmode+nres)
С
С
                                    (dimag(freq(i)),i=nmode+nres,2*(nmode+nres))
   *** The next 3 records are repeated (as a pair)
С
   *** The number of iterations is stored as ncumit.
C
С
   DSP
            cmp ncumit nmrow+5/ Calculated complex displacements
С
                          nmUsed+5 The first nmrow entries are the displacements
                                    in the same order as the original set of DOFs
C
С
                                    (see record AFTER ptrDOF). If DSPfmt=0, these
С
                                    are physical displacements, If DSPfmt!=0,
                                    these are the nmUsed modal coordinates instead
C
                                    of the nmrow entries.
С
С
С
                                    For the last five entries:
C
                                       Real part
                                                              Imag part
                                    1. frequency for these
                                                              frequency increment
C
С
                                    2. load step number
С
                                                              substep number
C
                                    3. cumulative iteration emrpm (MRPM cmd)
С
                                       number
С
                                    4. zero
                                                              zero
C
                                    5. scale factor
                                                              numdeflvs
С
                                         (cvs(i),i=1,nmrow/nmUsed),(freq,delf),
                                         (itime, itter), (ncumit, 0.0), (0.0, 0.0),
С
                                         (fscale, numdeflvs)
C
                                    Note: If, upon reading of this record, there
C
С
                                    is less than nmrow+5 items in the record,
С
                                    then only a selected set of nodes were
                                    output. Another record follows (integer, less
C
                                    than lenbac long) which contains the list of
С
С
                                    nodes for which DOF solutions are available.
```

```
c --- i 1 numdeflvs lvscal table scale factor IDs
c (ilvscID(i),i=1,numdeflvs)

c --- dp 1 numdeflvs lvscal table scale factor values
c (dlvscVal(i),i=1,numdeflvs)
```

1.5. Description of the Modal Results File

This section explains the content of the modal results file (jobname.mode).

1.5.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 9.

1.5.2. MODE File Format

```
*comdeck,fdmode
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
     ****** description of modal result file *******
     character*8 MODENM
     parameter
                (MODENM='mode
     character*8 MODENM_LEFT
     parameter (MODENM_LEFT='lmode
     *** NOTE: if this variable is changed in the future it should be
C
С
               updated in spdefines.h also for symbolic assembly (jrb)
                    MODEHDLEN
     integer
                    (MODEHDLEN=100)
     parameter
     LONGINT
                     modefpL
     integer
                     modebk, modeut
     LONGINT
                     modeLeftfpL
     integer
                     modeLeftbk, modeLeftut
     common /fdmode/ modefpL, modebk, modeut,
                     modeLeftfpL, modeLeftbk, modeLeftut
   ****** common variable descriptions *******
С
               file position on file mode
CO
CO
  modebk
                block number for file mode
   modeut
                file unit for file mode
CO
   modeLeftfpL file position on file .lmode
co modeLeftbk block number for file .lmode
   modeLeftut
               file unit for file .lmode
   See fddesc for documentation of how binary files are stored.
С
С
     ****** file format ******
C
        recid tells the identifier for this record. Not all records will have
C
             identifiers -- they are only indicated for those records whose
С
             record pointers are stored in the second file header.
        type tells what kind of information is stored in this record:
```

```
С
              i - integer
              dp - double precision
С
              cmp - complex
C
С
         nrec tells how many records of this description are found here
         lrec tells how long the records are (how many items are stored)
c recid
           type
                   nrec
                           lrec
                                     contents
            i
                    1
                           100
                                     standard ANSYS file header (see binhed8 for
C
                                     details of header contents)
С
            i
                    1
                           100
                                     .MODE FILE HEADER
C
С
                                                             0,
                                      fun09,
                                                                    nmode, numdof, < 5
С
                                                nmrow.
                                                         lenbac, nEnfGrp,
                                                                               negns,
С
                                      maxn,
                                                wfmax,
                                                                   kan,
                                                                              ldstep, < 15
С
                                     lumpms,
                                               extopt, SvCode,
                                     numitr, expbeg, expend,
                                                                   nspect, nSPdat, < 20
C
                                          0, ptrFRQ, kPerturb, ptrSHP, ptrLOD, < 25
C
                                     ptrNAR, ptrNARh, ptrDMP, nrkeyPert, nrigid, < 30
C
                                     ptrLPM, ptrSP1, ptrSHPh, ptrLODh, 0, < 35
0, ptrDMPh, ptrLPMh, ptrSP1h, ptrIRHS1, < 40
trIRHSh, 0, ptrRES, ptrRESh,Glblenbac, < 45
С
С
                                   ptrIRHSh,
C
                                  KeyStress, ptrELD, ptrELDh, ptrGBk, ptrGBkh, < 50
С
                                              nresi, ptrEf1, ptrEf1h,
С
                                    modlstp,
                                                                             sstif, < 55
                                    ptrFSTA, ptrEf2, ptrEf2h, ptrEf3h, < 60
C
                                                                   ptrHI, ptrKUNS,
                                  qrDampKey,cycMSUPkey,cycnmode,
С
                                   ptrKUNSh, mrestart, LPrestls, LPrestss, cpxmod, < 70
С
                                    keyLeft, cpxlv, ptrSCL, sparseLV, udfrqkey, < 75
С
С
                                  ptrUDFRQl,ptrUDFRQh,
                                                             0, 0,
                                                                                0, < 80
                                                   0,
                                                              0,
                                          0,
                                                                        0,
                                                                                   0, < 85
C
                                                                                   0, < 90
0, <100
С
                                          Ο,
                                                    0,
                                                               Ο,
                                                                         0,
С
                                          0,
                                                    0,
                                                               0,
                                                                         0,
                                    each item in header is described below:
С
С
                                     fun09 - unit number (mode file is 9)
                                     nmrow - number of rows/columns in matrices
С
                                              (maxn*numdof)
С
                                     nmode - number of modes extracted
C
С
                                     numdof - number of dof per node
                                     maxn - maximum node number (if extopt=3
С
С
                                              or 4, the actual number of nodes is
                                             referenced.)
С
С
                                     wfmax - maximum wavefront (does not apply
С
                                              if extopt=3 or 4)
                                     lenbac - number of nodes
C
                                    nEnfGrp - numbre of enforced group
neqns - number of equations on the .LN22
С
С
С
                                              file
                                     lumpms - lumped mass key
C
                                              = 0 - default matrix type
C
                                              = 1 - lumped
С
С
                                              (does not apply if extopt=3 or 4)
                                     extopt - mode extraction method
C
С
                                              = 3 - unsymmetric Lanczos
С
                                              = 4 - damped Lanczos
                                              = 6 - block Lanczos
C
                                              = 8 - SuperNode
С
                                              = 9 - PCG Lanczos
С
                                     SvCode - Solver assembly code path
C
С
                                              = 1 Symbolic assembly
                                            - analysis type
C
                                     kan
С
                                              = 1 - buckling
                                              = 2 - modal
С
                                     ldstep - load step number - also number of load vectors
C
                                     numitr - total number of cumulative
С
                                              iterations done during analysis
```

```
С
                                                (does not apply if extopt=3 or 4)
С
                                      expbeg - beginning of the frequency range of
                                                interest.
C
С
                                      expend - end of the frequency range of
С
                                                interest
                                      nspect - number of spectra; if -6, these are
C
                                                the 6 default unit spectra
С
                                      nSPdat - number of data items per spectrum
С
                                      ptrFRQ - pointer to the frequencies
C
C
                                    kPerturb - Linear Perturbation key
                                      ptrSHP - pointer to the mode shapes
C
С
                                                (eigenvectors)
                             ptrLOD, ptrLODh - pointer to the load vectors
C
                             ptrNAR, ptrNARh - pointer to the nodal averaged result records
C
                            ptrDMP, ptrDMPh - pointer to the modal damping matrix
С
С
                          ptrKUNS, ptrKUNSh - pointer to the modal stiffness
                                               matrix (unsymmetric part)
С
                                   nrkeyPert - nrkey setting of base analysis (Linear Perturbation)
С
                                      nrigid - number of rigid body modes
С
                                      ptrLPM - pointer to the diagonal mass vector
С
                                      ptrSP1 - pointer to the the spectrum data
C
                                  ptrIRHSl,h - pointer to imaginary part of RHS vector
C
                             ptrRES, ptrRESh - pointer to residual vectors
    Glblenbac - global number of nodes (for D-ANSYS)
С
C
                              ptrGBk,ptrGBkh - pointer to global nodal equivalence table
C
                                     modlstp - multiple load step key
С
С
                                       nresi - number of residual vectors in file
                                   KeyStress - key set if mode stresses on file
C
                             ptrELD,ptrELDh - pointer to element records
ptrEf1,ptrEf1h - pointer to enforced motion dof information
С
С
                                       sstif - key denoting prestress effects are included,
С
                                                this key is for internal usage only. SSTIF, on
C
                                                or off is controlled by NLGEOM on or off now.
C
                             ptrFSTA - pointer to fstacm data
ptrEf2,ptrEf2h - pointer to enforced motion modes
C
С
                              ptrEf3,ptrEf3h - pointer to enforced motion force
C
С
                                   qrDampKey - QR damped calcaulations key
                                  cycMSUPkey - mode file format is for subsequent cyclic MSUP
С
                                                (only base results on file)
C
С
                                    cycnmode - total number of cyclic modes extracted
                                                (sum of all harmonic indices)
С
                                       ptrHI - pointer to harmonic indices
C
                                    mrestart - key for modal restart (=0 none, =1 modal restart)
C
C
                                    LPrestls - restarted load step (from linear perturbation)
С
                                    LPrestss - restarted substep (from linear perturbation)
                                      cpxmod - key for complex frequencies/shapes (0=no 1=yes)
С
                                     keyLeft - key for LMODE writing (0=no 1=yes)
C
С
                                       cpxlv - key for RHS vector in complex form (0=no 1=yes)
С
                                                = 0 no (before version 17.0)
                                                = 1 yes (since version 17.0); ptrIRHSl,h is 0
C
                                      ptrSCL - pointer to cyclic mode scale factors
С
                                                (if scaled to unity, modnrm on modopt)
С
                                    sparseLV - key if load vectors have been sparsified
С
                                           0 - position not used
C
                                    udfrqkey - Key for format and writing of undamped frequency record
C
                        ptrUDFRQ1,ptrUDFRQh - pointer to undamped frequency record
С
                     1
                                      Degrees of freedom per node
C
                           numdof
                                      DOF reference numbers are:
С
          UX = 1, UY = 2, UZ = 3, ROTX= 4, ROTY= 5, ROTZ= 6, AX = 7, AY = 8
С
          AZ = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
C
           CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
С
          EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
                                       (curdof(i), i=1, numdof)
C
                     1
                            lenbac
                                      Nodal equivalence table
C
С
                                      This table equates the number used for
С
                                      storage to the actual node number.
                                       (Back(i), i=1, lenbac)
C
c GBK
                     1
                          Glblenbac Global nodal equivalence table
```

```
С
                                      This table equates the number used for
                                      storage to the actual global node number.
С
                                       (GlbBack(i), i=1, Glblenbac)
C
С
  FSTA
           dр
                     1
                           30
                                      fstacm.inc information (mass and MofI)
            i
                                      Signed harmonic index for each extracted frequency.
С
   ΗI
                     1
                           cycnmode
                                      Only present if cycMSUPkey=1.
С
C
   FRQ
           dp/cmp
                     1
                           nf
                                      Frequencies (eigenvalues).
                                      Frequencies are complex if cpxmod=1 or qrdampKey=1.
С
С
                                      If frequencies are real, numbers stored are
С
                                         the squares of the natural circular
                                         frequencies (w^**2, where w=radians/time).
С
                                         You can obtain the natural frequencies, f
С
С
                                         (in cycles/time), using the equation f=w/2pi
                                      If frequencies are complex, numbers stored are
С
                                         the natural frequencies (Hz)
С
С
                                      (freq(i), i=1, nf)
                                          nf = nmode+nresi
C
                                        if cycMSUPkey=1 then
С
                                          nf = cycnmode
C
   UDFRQ
                    1
                          nf
                                      Undamped Eigenvalues for a QRDAMP Modal Analysis.
С
             dp
                                      Record only exists when udfrqkey=1.
C
                                      Numbers are stored as the squares of the natural
С
С
                                      circular frequencies.
С
                                      This record is directly written to the mode file and might
                                      not be present in the .modesym file
С
                                      Mode shapes (eigenvectors). Mode shapes are
   SHP
           dp/cmp
                           nmrow
C
                     ns
                                      complex if cpxmod=1. The order corresponds
С
С
                                      to the nodal equivalence table
                                       (psi(i,j),i=1,nmrow)
С
С
                                          ns = nmode
                                       if cycMSUPkey=1 then
C
С
                                          ns = cycnmode
   RES
            dp
                                      Residual vectors
C
                   nresi
                           nmrow
                                      Load vectors in complex form (since version 17.0)
С
   LOD
           cmp
                  ldstep
                           nmrow
                                       (f(i), i=1, nmrow)
C
                                      Before version 17.0, records were (cpxlv=0):
С
С
                                          dp ldstep nmrow Load vectors
С
                                                             (f(i), i=1, nmrow)
С
                                      IRHS dp ldstep nmrow Imaginary Load vectors
                                                             (fimag(i), i=1, nmrow)
C
С
   LPM
            dp
                     1
                           nmrow
                                      Lumped mass vector. This record is present
                                      only if lumpms=1 and nmatrix=0. It is a
C
                                      vector containing the mass at each node in
С
С
                                      the system.
                                       (mass(i),i=1,nmrow)
C
   DMP
                                      Modal damping matrix. Each row of the matrix
C
            dp
                   nmrow
                           nmrow
C
                                      matrix is stored as a record.
С
                                       (ac(i,j),i=1,nmrow)
С
   KUNS
                                      Modal unsymmetric stiffness matrix. Each row of the
            dp
                   nmrow
                           nmrow
С
                                      matrix is stored as a record.
                                       (aku(i,j),i=1,nmrow)
С
                                      (groupID(i), i=1, nEnfGrp)
С
   EF1
           int
                     1
                          nEnfGrp
                                      (grpdof(i),i=1,nEnfGrp)
           int.
                     1
                          nEnfGrp
C
                          grpdof(i)
С
                 nEnfGrp
                                      dofIndx(i,j) i=1,grpdof(j)
                                      The above records contain information about each
С
                                      enforced motion group.
C
                                      Enforced static modes
С
  EF2
                nEnfGrp
                           nmrow
c EF3
            dр
               nEnfGrp
                           nmrow
                                      Enforced forced vector
```

```
c for each spectrum (|nspect| records):
                        nmode+nresi Participation factors for this spectra
  SP1
            dр
C
                    1
С
            dр
                    1
                        nmode+nresi Mode coefficients for this spectra
С
   ---
            ďρ
                    1
                        nmode+nresi Modal damping values
                    1
                         ndsvc*
                                       (*) see svcom.inc
C
            ďρ
            int
                    1
                         nisvc*
С
С
            dр
                    1
                          20
                                     misc. spectra data
С
   ELD
            int
                    1
                          15
                                     nelm, maskl, nItems, ptrELM, ptrERS,
                                   ptrCER1,ptrCERh, ptrESL1,ptrESLh,
                                                                        nRF,
                                                                                 (10)
C
С
                                    ptrRFl, ptrRFh, PrecKey, maskh,
                                    each item in header is described below:
C
С
                                      nelm - number of elements
                                   maskl,h - output mask (OUTRES)
C
                                    nItems - number of element records (7, VOL
С
С
                                             not included)
                                    ptrELM - pointer to element equivalence table
C
                                   ptrERS - pointer to element record sizes
C
                                 ptrCER1,h - pointer to constant element records
C
                                 ptrESLl,h - pointer to element index
С
                                       nRF - number of reaction forces
C
                                  ptrRFl,h - pointer to reaction forces
C
                                   PrecKey - 0, double precision 1, single
С
С
                       above pointers are relative to ptrELD
                                   Total size of each element record (LONGINT)
C
            int
                        2*nItems
                    1
                          nelm
                                    Element equivalence table
   ELM
            int
C
                                    This table equates the order number used to
C
                                     the actual element number
C
                                    Sizes of the nItem element results sets for
С
   ERS
            int
                  nItems nelm
                                     each element
C
С
   CER
            int
                    1
                          5
                                  ptrVOL, ptrEPT, ptrEUL,
                                                                0,
                                                                         0
                       above pointers are relative to ptrCER
C
      constant element records (do not vary by mode):
С
    VOL
                    1 nelm*1
                                    Element volume
C
            dp
                    1 nelm*size
                                     Element structural nodal temperatures
С
    EPT
            dp
C
    EUL
            dр
                    1 nelm*size
                                    Element Euler angles
    ESL
            int
                    1
                          10
                                   ptrENS, ptrEEL, ptrEMS, ptrENF, ptrENG,
C
                                   ptrENSh,ptrEELh,ptrEMSh,ptrENFh,ptrENGh
C
С
                       above pointers are relative to ptrESL
     non-constant element records (do vary by mode):
C
                 nelm nmode*size
                                    Element nodal component stresses
С
            dр
С
    EEL
            dр
                 nelm nmode*size
                                    Element nodal component elastic strains
    EMS
            dp
                 nelm nmode*size
                                    Element summable miscellaneous data
C
    ENF
                 nelm nmode*size
                                    Element nodal forces
С
            dp
                 nelm nmode*3
    ENG
            dp
                                     Element energies
C
      see fdresu.inc for more information on the element results
C
   NAR
            int
                                      nnod, ptrNCTl, ptrNCTh, ptrNSTl, ptrNSTh,
С
С
                                      ptrNELl, ptrNELh
С
                                    each item in header is described below:
                                     nnod - number of nodes
C
                                 ptrNCTl,h - pointer to node contributions
C
                                 ptrNSTl,h - pointer to NAR stress record
C
С
                                 ptrNELl,h - pointer to NAR elastic strain record
C
                       above pointers are relative to ptrNAR
      NAR records (do vary by mode):
C
          i nnod 2
                                   Node Contributions
```

```
c NST dp nnod nmode*6 Nodal component stresses
c NEL dp nnod nmode*7 Nodal component elastic strains
c see fdresu.inc for more information on the NAR records
```

1.6. Description of the Element Matrices File

This section explains the content of the element matrices file (jobname.emat).

1.6.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 2.

1.6.2. EMAT File Format

```
*comdeck,fdemat
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
С
      ****** description of element matrix file *******
C
      character*8 EMATNM
     parameter (EMATNM='emat
                   (EMATHDLEN=80)
     parameter
     LONGINT
                     ematfpL, ematfp
      integer
                     ematbk, ematut, maxldset
      common /fdemat/ ematfpL, ematbk, ematut, maxldset
      equivalence (ematfp,ematfpL)
    ******* common variable descriptions *******
С
  ematfpL
                file position on file emat
CO
                block number for file emat
    ematbk
CO
    ematut
                file unit for file emat
C
    See fddesc for documentation of how binary files are stored.
С
      ****** file format ******
C
С
        recid tells the identifier for this record. Not all records will have
             identifiers -- they are only indicated for those records whose
C
              record pointers are stored in the second file header.
C
         type tells what kind of information is stored in this record:
C
              i - integer
С
             dp - double precision
С
             cmp - complex
C
        nrec tells how many records of this description are found here
C
         lrec tells how long the records are (how many items are stored)
c recid
           type
                   nrec
                          lrec
                                   contents
                   1
                          100
                                   standard ANSYS file header (see binhed8 for
С
                                   details of header contents)
                           80
                                 .EMAT FILE HEADER
```

```
С
                                               nume, numdof,
                                                                   lenu, lenbac,
С
                                      fun02,
                                      maxn, nlgeEMA, sstEMAT,
                                                                   0, lumpm,
C
                                                        kycd, kygss,
С
                                      kygst, kygm,
                                                                            kygaf,
С
                                      kygrf,
                                                   0,Glblenbac, ptrGBkl, ptrGBkh,
                                                   0, 0, ptrBITh, ptrEHDh,
C
                                    ptrElmh.
                                     ptrIDXh, numCE, maxLeng, ptrCEl, ptrCEh, ptrDOF, ptrBAC, ptrELMl, 0, 0,
                                    ptrIDXh,
С
С
                                    ptrBITl, ptrEHDl, ptrIDXl, ptrendH, ptrendL,
C
С
                                     nldstp, maxldset, ptrLSI1, prtLSIh,
                                                   0,
                                                             0,
                                                                      0,
                                          0,
                                                                                0,
C
                                          0,
                                                   0,
С
                                                              Ο,
                                                                       0,
С
                                          0,
                                                   0,
                                                             0,
                                                                       0,
                                                                                0,
                                          0,
                                                   0,
                                                              0,
                                                                       0,
                                                                                0,
C
                                          0,
C
                                                   0,
                                                              0,
                                                                       0,
                                          0,
С
                                                   0,
                                                              0,
                                                                       Ο,
                                                                                0,
                                          0,
                                                   0,
                                                              0,
                                                                       0,
С
С
                                  each item in header is described below:
                                     fun02 - unit number (emat file is 2)
C
                                     nume - number of elements
C
                                     numdof - number of dofs per node
С
                                     lenu - total DOFs of model
C
                                     lenbac - number of nodes
C
                                     maxn - maximum node number
С
                                     nlgeEMA = 0 - nlgeom is OFF the time this Emat file is created
С
                                                1 - nlgeom is ON the time this Emat file is created
C
                                     sstEMAT = 0 - sstif key is OFF the time this Emat file is created
С
                                                {\bf 1} - sstif key is ON the time this {\tt Emat} file is created
С
                                                    this key is for internal use only
С
                                     lumpm - lumped mass key
C
                                              = 0 - default matrix type
C
                                              = 1 - lumped
С
                                     kygst - global stiffness matrix calculate
С
С
                                              kev
С
                                              = 0 - do not calculate
С
                                              = 1 - calculate
                                            - global mass matrix calculate key
C
                                     kyqm
С
                                              = 0 - do not calculate
                                              = 1 - calculate
С
                                            - global damping matrix calculate key
С
                                     kycd
                                              = 0 - do not calculate
C
С
                                              = 1 - calculate
С
                                     kygss - global stress stiffening matrix
С
                                              calculate key
                                              = 0 - do not calculate
С
С
                                              = 1 - calculate
С
                                     kygaf - global applied force vector
                                              calculate key
C
                                              = 0 - do not calculate
С
                                              = 1 - calculate
С
С
                                     kygrf - global restoring force vector
                                              calculate key (Newton-Raphson only)
C
                                              = 0 - do not calculate
C
С
                                              = 1 - calculate
                                            - position not used
С
                                     Glblenbac - global global number of nodes (== lenbac unless using
C
С
                                                 Distributed Ansys)
С
                                     ptrGBkl- low pointer to global nodal equivalence table
                                     ptrGBkh- high pointer to global nodal equivalence table
C
С
                                     ptrELMh- high pointer to element equivalence table
                                     ptrBITh- high pointer to dof bits
С
                                     ptrEHDh- high pointer to the start of the
C
С
                                              element matrices
                                     ptrIDXh- high pointer to element matrices
C
С
                                              index table
                                     numCE - number of internal CEs
С
                                     maxLeng- maximum length of any internal CE
C
                                     ptrCEl - low pointer to internal CE list
С
С
                                     ptrCEh - high pointer to internal CE list
```

```
С
                                    ptrDOF - pointer to degrees of freedom per
С
                                             node used in model
                                    ptrBAC - pointer to nodal equivalence table
C
С
                                    ptrELM1- Low pointer to element equivalence
                                             table
C
                                    ptrBITl- Low pointer to dof bits
С
                                    ptrEHDl- Low pointer to the start of the
С
                                             element matrices
C
С
                                    ptrIDX1- Low pointer to element matrices
                                             index table
С
С
                                    ptrendH- High pointer to end of file
                                    ptrendL- Low pointer to end of file
C
С
С
                                    nldstp - number element load vector set
                                    maxldset- max eload vector set
С
                                    ptrLSIl - location of Load Step Index
С
С
                                    prtLSIh -
        Note: the analysis type sets the global calculate keys.
C
           dр
                    1
                            20
                                    Time information
С
С
                                    timval, timinc, frqval, timbeg, timend,
C
                                               0.0,
                                                       0.0,
С
                                       0.0,
                                                               0.0,
                                                       0.0,
                                               0.0,
                                                               0.0,
                                                                       0.0,
С
                                       0.0,
C
                                       0.0,
                                               0.0,
                                                       0.0,
                                                               0.0,
                                                                       0.0,
                                   each item is described below:
С
                                    timval - the current time
C
                                    timinc - the time increment
C
                                    frqval - the current frequency (from a
С
С
                                             harmonic analysis)
                                    timbeg - the start time for the analysis
С
С
                                    timend - the end time for the analysis
С
                                    0.0
                                           - position not used
                                    0.0
                                           - position not used
С
С
                                    0.0
                                           - position not used
                                           - position not used
                                    0.0
С
                                           - position not used
                                    0.0
С
                                    0.0
                                           - position not used
С
С
                                    0.0
                                           - position not used
С
                                    0.0
                                           - position not used
С
                                    0.0
                                           - position not used
                                           - position not used
                                    0.0
С
С
                                    0.0
                                           - position not used
С
                                    0.0
                                           - position not used
                                    0.0
C
                                           - position not used
                                           position not usedposition not used
                                    0.0
С
С
                                    0.0
                         numdof
                                    Degrees of freedom per node
С
                                    DOF reference numbers are:
C
          С
С
          CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
C
          EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
С
                                     (curdof(i), i=1, numdof)
С
   BAC
                    1
                         lenbac
                                    Nodal equivalence table. This table equates
                                    the number used for storage to the actual
С
                                    node number
С
С
                                     (Back(i), i=1, lenbac)
С
   ELM
                    1
                                    Element equivalence table. The ANSYS program
                          nume
С
                                    stores all element data in the numerical
                                    order that the SOLUTION processor solves the
С
                                    elements. This table equates the order
С
С
                                    number used to the actual element number
```

```
(Order(i), i=1, nume)
С
   GBK
            i
                    1 Glblenbac
                                     Global nodal equivalence table. This
C
С
                                     table equates the number used for storage
С
                                     to the actual node number. Only written
                                     by the master process in Distributed Ansys
C
                                      (GlbBack(i), i=1, Glblenbac)
С
                    1
                          lenu
                                     Bits set at a DOF table. This record
C
   BIT
            i
C
                                     has bits for constraints, forces, etc.
                                      (DofBits(i), i=1, lenu) (added at 10.0)
С
С
   IDX
            i
                    1
                           2*nume
                                     Element index table. This record specifies
                                     the file location for the beginning of the
C
                                     data for each element.
С
С
                                      (index(i),i=1,nume) Low part of pointer
                                      (index(i),i=1,nume) High part of pointer
С
c The records at the end of the file store element information and get written
c as a set for each element(nume sets of these records will appear on the file
c at this point) ptrEHD indicates the beginning of the element data.
c If substructure matrices are written to the EMAT file, they are written in a
  different format than is shown here. This alternate format is not documented
  at this time, as it is likely to change in the future.
   EHD
            i
                    1
С
                           10
                                     Element matrix header
С
                                               mkey,
                                                       dkey, sskey, akey,
C
                                      stkev,
                                              ikey, kckey,
C
                                    each item in header is described below:
C
                                     stkey - stiffness matrix key
C
С
                                              = 0 - matrix not present
С
                                              = 1 - matrix present
                                            - mass matrix key
С
                                     mkev
                                              = 0 - matirx not present
С
                                              = 1 - matrix present
С
                                     dkey
                                            - damping matrix key
С
                                              = 0 - matrix not present
С
С
                                              = 1 - matrix present
С
                                     sskey
                                            - stress stiffening matrix key
С
                                              = 0 - matrix not present
                                              = 1 - matrix present
С
С
                                     akev
                                            - applied load vector key
С
                                              = 0 - vector not used
                                              = 1 - vector used
C
                                     nrkey - newton-raphson(restoring) load
С
С
                                              vector key (for nonlinear analyses)
С
                                              = 0 - vector not used
                                              = 1 - vector used
С
                                     ikev
                                            - imaginary load vector key
C
С
                                                (for complex analyses)
С
                                              = 0 - vector not used
                                              = 1 - vector used
C
С
                                     kckey
                                              = 0 or 1 or 2 position for internal use or not in use
С
                                              = 3 complex number stiffness matrix key
C
                                            - position not used
С
С
                                     nmrow
                                           - numbers/columns in matrices. If the
                                              number is negative, the matrices
C
С
                                              will be written in upper triangular
                                              form.
С
C
                    1
                         nmrow
                                     DOF index table. This record specifies the
                                     DOF locations of this element matrix in
С
                                     relation to the global matrix. The index is
С
С
                                     calculated as (N-1)*NUMDOF+DOF, where N is
```

```
С
                                     the position number of the node in the nodal
                                     equivalence table and DOF is the DOF
С
                                     reference number given above
C
С
           dр
                 varies varies
                                     Element matrices. This record is repeated
                                     for each stiffness, mass, damping, stress stiffening
C
                                     and complex stiffness matrice. If the matrix is
С
                                     diagonal, the length of the records will be
С
                                     nmrow. If the matrix is unsymmetric, the
C
C
                                     length of the records will be nmrow*nmrow.
                                     If the matrix is symmetric, only the upper
С
С
                                     triangular terms are written and the length
С
                                     of the records will be (nmrow)*(nmrow+1)/2.
                                     Element force vectors. This record contains
С
           dр
                          2*nmrow
С
                                     both the applied force vector and the
                                     (restoring or imaginary) load vector.
С
С
С
       ********* Internal CE information ***********
C
       The following records repeat numCE times... one for each internal
C
       CE created during solution... these are stored here for the
С
       usage of a prestressed modal analysis such as the linear perturbation analysis
С
С
                                     First part is the CE number, the second part is
   CE
           i
С
                         numCE
                                     the number of terms in this internal CE, and
С
                                     the third part is the external element number
С
C
                                     of the element that created this internal CE
С
                                     integer info (list of node*32 + dof)
С
           i
                 nTerms
                         numCE
C
                                     dp info (list of coefficients including constant term)
С
   ___
          dp
                 nTerms
                         numCE
С
C
C
    kygst
                 global stiffness matrix calculate key
                 global mass matrix calculate key
С
    kygm
С
    kygd
                 global damping matrix calculate key
С
                 global stress stiffening matrix calculate key
    kygss
    kygaf
                 global applied force matrix calculate key
C
                 global restoring force matrix calculate key
С
    kygrf
С
C
С
С
  Additional element records stored to support topo optimization
С
   with multiple load steps. Currently only element load vectors are needed,
   but keep the same format of the original element matrices/load vectors
   for possible future need.
C
С
С
  header(41) - header(44) are all 0 unless it's topo opitimization analysis
C
                          2*maxldset Load index table. This record specifies
С
С
                                      the file location for the beginning of the
С
                                      load steps after load step 2. In each load
                                      steps, we have the following two records,
С
                                      TPIDX and element force vectors
С
C
С
   TPIDX
            i
                    1
                          2*nume
                                      Topo index table. This record specifies
                                      the file location for the beginning of the
C
С
                                      data for each element record, EHD.
С
                                       (index(i),i=1,nume) Low part of pointer
                                       (index(i),i=1,nume) High part of pointer
С
С
                                      same as IDX above
С
    EHD
                                      Element matrix header
С
            i
                    1
                          10
С
                                      same as EHD above
С
С
                    1
                          nmrow
                                      DOF index table
С
                                      Element force vectors. This record contains
С
           dp
                    1
                          2*nmrow
                                      both the applied force vector and the
С
                                      (restoring or imaginary) load vector.
C
```

1.7. Description of the Substructure Matrices File

This section explains the contents of the substructure matrices file (jobname.sub).

1.7.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 8.

1.7.2. SUB File Format

```
*comdeck,fdsub
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc
      ******* description of substructure matrix file *******
      character*8 SUBNM
     parameter (SUBNM='sub
     integer
                    SUBHDLEN
     parameter
                  (SUBHDLEN=80)
     LONGINT
                    subfpL, lenSubL
                    subbk, subut
      common /fdsub/ subfpL, lenSubL, subbk, subut
c write: matout
c read:
    ****** common variable descriptions ********
С
              file position on file sub
  subfpL
CO
  subbk
               block number for file sub
CO
CO
  subut
               file unit for file sub
   lenSubL
               length of sub file (saved for slvdta.F)
    See fddesc for documentation of how binary files are stored.
C
C
     ****** file format *******
С
     recid tells the identifier for this record. Not all records will have
С
          identifiers -- they are only indicated for those records whose
С
          record pointers are stored in the second file header.
C
С
      type tells what kind of information is stored in this record:
          i - integer
C
          dp - double precision
С
          cmp - complex
С
     nrec tells how many records of this description are found here
      lrec tells how long the records are (how many items are stored)
c recid
          type
                  nrec
                          lrec
                                   contents
           i
                   1
                          100
                                   standard ANSYS file header (see binhed8
C
                                   for details of header contents)
С
  HED
                           80
                                   .SUB FILE HEADER (FULL MATRICES)
С
                   1
C
```

```
8, nmrow, nmatrx, nedge, numdof, maxn, wfmax, lenbac, nnod, kunsym, 10
С
С
                                      kstf, kmass, kdamp, kss, nvect,
C
                                    nWorkL, lenU1, sesort, lenlst,ptrLodL,
С
С
                                    ntrans, ptrMtx, ptrXFM, ptrHED, name1,
                                   name2, ptrCG, 0, name3, name4, ptrDOF, ptrDST, ptrBAC, ptrTIT, ptrNOD, ptrXYZ, ptrEDG, ptrGDF, thsubs, ptrPOS,
C
С
С
                                    ptrORG, stfmax, ptrLodH, nmodes, keydim,
C
                                 cmsMethod, name5, name6, name7, name8,
nvnodes,ptrCTXM, nWorkH, 0,ptrTVAL,
C
С
                                  gyroDamp,kstress,nStartVN,ptrEndL,ptrEndH,
С
                              ptrimsSEdat,ptrdmsSEdat,units, 0,
                                                                           0,
                                         0,
                                                         0,
                                                                    0,
                                                                            0,
                                                 0,
                                                                                 70
C
                                                                   0,
                                         0,
                                                  0,
                                                            0,
C
                                                  0,
                                                           0,
                                                                    0,
                                                                            Ω
С
                                         0,
                                                                                 80
                                       .SUB FILE HEADER (SPARSE MATRICES)
С
   HED
             i
                    1
                              80
C
                                         9, nEqn, nmatrx, ndege, numdof,
C
                                      maxn, wfmax, lenbac, nnod, kunsym,
C
С
                                      kstf, kmass, kdamp,
                                                                , nvect,
                                    nTermL,
                                              , , lenlst,ptrLodL,
С
                                    ntrans,ptrMtxL, ptrXFM, ptrHED, name1,
C
                                    name2, ptrCG, , name3, name4,
С
                                    ptrDOF, , ptrBAC, ptrTIT, ptrNOD,
С
C
                                    ptrXYZ, ptrEdg, ptrGDF, thsubs, ,
                                          , stfmax,ptrLodH, , keydim,
, name5, name6, name7, name8,
С
С
                                          ,ptrCTXM, nTermH,ptrMtxH,ptrColL,
С
                                   ptrColH,ptrCofL,ptrCofH,ptrEndL,ptrEndH,
C
С
С
                                                                                70
C
С
                                                                                 80
                                   each item in header is described below:
C
                                    fun08 - unit number (full sub file is 8)
С
                                             (sparse substructure file is 9)
C
                                    nmrow - number of rows in matrices (also
C
                                            number of dofs in substructure)
C
                                    nmatrx - number of matrices on file
С
                                    nedge - number of edges for outline
numdof - number of dofs per node
С
C
С
                                    maxn - maximum node number of complete
С
                                            model presently in database
                                    wfmax - maximum wavefront of substruct.
C
                                             during generation pass
С
                                    lenbac - number of nodes defining
С
                                             substructure during the
C
                                            generation pass
C
                                    nnod - number of unique nodes in the
C
С
                                              substructure having DOFs, and
С
                                              which define this substructure
                                              during the use pass. Also, the
C
                                             number of nodes having master
С
С
                                             DOFs.
C
                                    kunsym - unsymmetric matrix key
                                             = 0 - symmetric
= 1 - unsymmetric
С
С
                                          - stiffness matrix present key
C
                                    kst.f
                                             = 0 - matrix is not on file
C
                                             = 1 - matrix is on file
C
С
                                    kmass - mass matrix present key
С
                                             = 0 - matrix is not on file
                                              = 1 - matrix is on file
C
                                              =-1 - Lumped mass vestor (Sparse only)
С
                                    kdamp - damping matrix present key
```

```
= 0 - matrix is not on file
С
                                            = 1 - matrix is on file
С
                                          - stress stiffening matrx present
С
                                  kss
С
                                            = 0 - matrix is not on file
С
                                           = 1 - matrix is on file
                                  nvect - number of load vectors
C
                                            (at least 1 is required)
С
                                  nWorkL,H - BCS workspace length (only for
С
C
                                              bacsub)
C
                                  nTermL,H - Number of terms in sparse matrix
                                  lenU1 - length of intermediate transformation
C
С
                                            vector
C
                                  sesort - DOF set sort key
                                            = 0 - numbers are not sorted
C
                                            = 1 - numbers are sorted in
C
С
                                                  ascending order
                                  len1st - maximum length of DOF set for
С
                                            this substructure (maxn*numdof)
С
                                  ptrLod - pointer to the start of the load
С
                                           vectors (see also ptrLodh)
С
                                  ntrans - transformed key
C
                                            = 0 - substructure has not been
C
С
                                                  transformed
                                            > 0 - substructure copied
C
                                                  from another substructure,
C
                                                  via either SESSYM or SETRAN
С
                                  ptrMtxL,H - pointer to the start of the
С
C
                                               substructure matrices (iDiagL for
                                               sparse matrices)
С
                                  ptrXFM - pointer to the substructure
C
                                           transformations
С
                                  ptrHED - pointer to the SUB file header
C
C
                                  namel - first four characters of the
                                            substructure file name, in
С
С
                                            integer form
                                  name2 - second four characters of the
C
С
                                            substructure file name, in
С
                                           integer form
                                  name3 - third four characters of the
C
С
                                            substructure file name, in
                                           integer form
С
                                  name4 - fourth four characters of the
С
                                           substructure file name, in
C
C
                                            integer form
                                  ptrDOF - pointer to the DOF/node list
С
С
                                  ptrDST - pointer to the local DOF set
                                  ptrBAC - pointer to the nodes comprising
С
С
                                           the substructure
                                  ptrTIT - pointer to the title
С
                                  ptrNOD - pointer to the unique nodes
C
                                           defining the substructure
С
                                  ptrXYZ - pointer to the coordinates of the
С
С
                                           unique nodes
                                  ptrEDG - pointer to the substructure edges
C
                                  ptrGDF - pointer to the global DOF set
C
                                  {\tt ptrCG}\, - pointer to the element mass information thsubs - element type key
С
С
                                            = 0 - structural
C
                                            = 1 - 1st order non-structural
С
С
                                                  (generally from thermal)
                                           = 2 - 2nd order non-structural
C
С
                                  ptrPOS - pointer to the sorted substructure
                                           DOF set to the original
С
                                  ptrORG - pointer to the DOF set of the model
C
С
                                           during the generation pass
                                  stfmax - maximum diagonal stiffness term
C
С
                                            (packed into an integer)
С
                                  ptrLodh- High 32 bits of 64 bit pointer
                                  nmodes - number of modes used to generate
C
С
                                            CMS s.e.
                                  keydim - dimensionality key
```

```
= 1 - axisymmetric
С
                                           = 2 - 2-D
С
                                            = 3 - 3-D
C
                               cmsMethod - component mode synthesis method
С
С
                                  name5 - fifth four characters of the
                                           substructure file name, in integer
C
С
                                  name6 - sixth four characters of the
С
                                           substructure file name, in integer
C
С
                                           form
                                  name7 - seventh four characters of the
С
С
                                           substructure file name, in integer
С
                                  name8 - eighth four characters of the
C
                                           substructure file name, in integer
С
С
                                           form
                                 nvnodes - number of virtual nodes that contain
С
                                           the modal coordinates
С
С
                                gyroDamp - gyroscopic damping matrix key
                                           = 0 - damping matrix is
С
                                                 not skew-symmetric if present
C
                                            = 1 - damping matrix is
C
С
                                                  skew-symmetric, due to
                                                  gyroscopic effect
С
                                 kStress - = 1 if modal element results are
C
                                           on the .cms file
С
                                nStartVN - starting number of the virtual nodes
С
C
                                           if defined with CMSOPT command
                                 ptrCTXM - coordinate transformation
С
                               ptrColL,H - pointer to the iCol sparse matrix
С
С
                                             arrav
                               ptrCofL,H - pointer to the of the
C
                                             sparse matrix Sk(1:nTerm),
C
                                             Sm(1:nTermL),Sc(1:nTermL),
С
С
                                             Ss(1:nTermL) Each matrix is a
                                             single large record
C
С
                               ptrEndL,H - Next location after end of file
С
                             ptrimsSEdat - pointer to integer record for stage data
                             {\tt ptrdmsSEdat\ -\ pointer\ to\ double\ precision\ records\ for\ stage\ data}
С
                                  units - unit system used
С
                                           =-1 - no /UNITS specification
С
                                           = 0 - user defined units
С
                                            = 1 - SI
C
                                           = 2 - CSG
С
                                            = 3 - U.S. Customary, using feet
С
С
                                            = 4 - U.S. Customary, using inches
                                            = 5 - MKS
C
С
                                            = 6 - MPA
C
                                            = 7 - uMKS
                                note: name1/2/3/4/5/6/7/8 are the
С
                                      inexc4 representation of the
С
С
                                      32 character filename.
                                      name1/2/5/6/7/8 will be "0"
С
                                      for pre rev 5.2 files - cwa
C
    Note: If ntrans > 0, records from position ptrDOF to ptrGDF will be
C
          identical to the data for the copied substructure.
C
                    1
                                  Substructure transformations (5*25 double
  XFM
            ďρ
                            125
C
С
                                  precisions). This record has meaning only
                                  if ntrans > 0. You can define up to five
С
                                  levels of transformations, with 25 variables
C
С
                                  in each level. Up to the first seven
                                  variables are used as follows:
С
С
С
                                  If the substructure was transferred (via the
                                   SETRAN command):
C
                                  1st variable - 1.0
С
                                  2nd variable - nodal increment
```

```
3rd variable - reference number of
С
С
                                    coordinate system where substructure will
                                    be transferred
C
С
                                   4th variable - reference number of
С
                                    coordinate system where substructure is
                                    presently defined
C
                                   5th variable - x coordinate increment
6th variable - y coordinate increment
С
С
                                   7th variable - z coordinate increment
C
                                   If the substructure used symmetry (via the
C
С
                                    SESYMM command):
С
                                   1st variable - 2.0
                                   2nd variable - nodal increment
C
                                   3rd variable - number of coordinate
С
С
                                    component to be used in operation
                                    = 1 - x coordinate
С
                                    = 2 - y coordinate
= 3 - z coordinate
С
С
С
                                   4th variable - reference number of
С
                                    coordinate system to be used for symmetry
                                    operation
C
С
   CTXM
            dр
                     1
                            250
                                   Substructure transformations
   DOF
                     1
                           numdof
                                      Degrees of freedom per node (Global)
C
                                       (curdof(i),i=1,numdof)
С
С
                                      DOF reference numbers are:
C
     UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8
         = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
С
     CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
С
     EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
            i
C
   DST
                     1
                           nmrow This record contains degrees of freedom for
                                   this substructure of the unique nodes, as
С
С
                                   used with this substructure, in ascending
                                   order. This index is calculated as
С
С
                                   (N-1)*numdof+DOF, where N is the node number
С
                                   and DOF is the DOF reference number given
                                   above
С
C
                                    (lsort(i),i=1,nmrow)
   POS
                     1
                                   This record stores the positions of the
С
                           nmrow
                                   local DOF set in relation to the generated
С
С
                                   DOF set. (lposit(i),i=1,nmrow)
С
   ORG
            i
                     1
                           nmrow
                                   DOF set of the model as defined during the
                                   generation pass. This index is calculated as
C
С
                                   (N-1)*NUMDOF+DOF, where N is the position
С
                                   number of the node in the nodal equivalence
С
                                   table and DOF is the DOF reference number
                                   given above
С
С
                                    (lorig(i),i=1,nmrow)
                           lenbac This group describes nodes that defined the
С
   BAC
                                   substructure during the generation pass of
C
                                   the analysis. Nodal data is stored in arrays
С
                                   equal to the number of used or referenced
С
                                   nodes. This table equates the number used
C
С
                                   for storage to the actual node number.
С
                                    (Back(i), i=1, lenbac)
С
   TIT
                     1
                              20
                                   Substructure title (converted to integers -
С
                                   see inexc4)
С
   NOD
                     1
                            nnod This record describes unique nodes defining
                                   the substructure for the use pass of the
С
С
                                   analysis. These are also the nodes having
С
                                   master degrees of freedom.
                                    (node(i), i=1, nnod)
C
c XYZ
           dр
                   nnod
                             6
                                   This record describes the coordinates of a
```

```
C
                                  unique node, in the order X, Y, Z, THXY,
С
                                  THYZ, and THZX. Nodal order corresponds to
                                  that of the node list given above
C
                                   (xyzang(j,i),j=1,6)
С
                  nedge
                             6
                                  This record contains beginning and ending
C
   EDG
           ďρ
                                  locations (X1,Y1,Z1,X2,Y2,Z2 coordinates)
С
С
                                  a straight line comprising an edge of the
                                  substructure.
C
          LONG
                                  This record describes global degrees of
С
   GDF
                     1
                           nmrow
С
                                  freedom of the unique nodes in ascending
С
                                  order, as used during the analysis use pass.
                                  This index is calculated as (N-1)*32+DOF,
С
С
                                  where N is the node number and DOF is the
С
                                  DOF reference number given above
С
                                   (l(i), i=1, nmrow) (sorted)
                                  (Made LONGINT in rev 14.0)
С
           dр
                                  Totmass, CGx, CGy, CGz, mm11, mm22, mm33, mm12, mm23,
C
   CG
                                  mm13. And three precise mass matrices: MASSTR,
C
                                  MASSROSP, and MASSTRRO. Each has 3*3 doubles.
С
   TVAL
                     1
                           1000
                                  current time value corresponds to each load step
C
C
                                  (substructuring analysis only)
С
   IMSSE
                     1
                            17
                                  integer record for stage data (see multiStage.inc):
                                  msNsector, msHindex, msNumoff,
C
                                  msCECYC(i)(i=1,2),msNodElm(i)(i=1,12)
С
                             6
                                  double precision record for stage data (see multiStage.inc):
   DMSSE
           ďρ
C
                                  msBox(i)(i=1,6)
C
c The substructure matrices are written at this position in the file. One row
  of each matrix is written to the file at a time. i.e. the first row of each
   matrix is written, then the second row of each matrix, etc. this pattern
C
   continues until all nmrow rows of each matrix have been written to the file.
   MAT
                                   Row of the stiffness matrix, if nmatrx > 0.
C
                    1
                           nmrow
С
                                    (ak(i,j),i=1,nmrow)
С
            dp
                           nmrow
                                   Row of the mass matrix, if nmatrx > 1.
                                    (am(i,j),i=1,nmrow)
C
                                   Row of the damping matrix, if nmatrx > 2.
С
                     1
                           nmrow
С
                                    (ac(i,j),i=1,nmrow)
С
                                   Row of the stress stiffening matrix, if
                           nmrow
С
                                   nmatrx > 3.
                                    (gs(i,j),i=1,nmrow)
C
C
  LOD
                           nmrow
                                   This record contains the load vectors.
                                    (f(i), i=1, nmrow)
C
```

1.8. Description of the Component Mode Synthesis Matrices (CMS) File

This section explains the contents of the CMS matrices file (jobname.cms).

1.8.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 8.

1.8.2. CMS File Format

```
*comdeck,fdcms
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
      ******* description of CMS (component mode synthesis) transformation file ********
      character*8 CMSNM
     parameter (CMSNM='cms
      LONGINT
                     cmsfpL, cmsfp
                     cmsbk, cmsut
     integer
      common /fdcms/ cmsfpL, cmsbk, cmsut
      equivalence (cmsfp,cmsfpL)
    ****** common variable descriptions *******
С
               file position on file cms
co cmsfp
               block number for file cms
  cmsbk
CO
  cmsut
               file unit for file cms
    See fddesc for documentation of how binary files are stored.
С
C
      ****** file format *******
С
С
         recid tells the identifier for this record. Not all records will have
              identifiers -- they are only indicated for those records whose
С
             record pointers are stored in the second file header.
С
         type tells what kind of information is stored in this record:
C
             i - integer
C
             dp - double precision
С
             cmp - complex
С
         nrec tells how many records of this description are found here
С
         lrec tells how long the records are (how many items are stored)
c recid
          type
                  nrec
                          lrec
                                   contents
С
                   1
                          100
                                   standard ANSYS file header (see binhed8 for
                                   details of header contents)
C
С
   ___
           i
                   1
                           40
                                   .CMS FILE HEADER
C
                                  fun45,
                                           neqn, nirfm, nnorm, ncstm,
С
                                  nrsdm, cmsMeth, kStress, lenbac, numdof,
С
                                            0,
                                cmsMixF,
                                                    0,
                                                                0,
                                                                          0,
C
С
                                      0,
                                               0,
                                                        0,
                                                                 0,
                                                                          0,
                                      0,
                                               0,
                                                        0,
                                                                0,
C
                                               0,
                                                        0, ptrNAR1, ptrNARh,
С
                                      0,
                                ptrIRF1, ptrNOR1, ptrCST1, ptrRSD1, ptrIRFh,
С
C
                                ptrNORh, ptrCSTh, ptrRSDh, ptrELDl, ptrELDh
                                  each item in header is described below:
С
                                   fun45 - unit number
C
С
                                    neqn - number of equations (DOF)
                                   nirfm - number of inertia relief modes
С
                                   nnorm - number of normal modes
С
С
                                   ncstm
                                          - number of constraint modes
                                          > 0 available in file
C
С
                                          < 0 not available in file
С
                                   nrsdm - number of residual modes
С
                                 cmsMeth - CMS method key
                                              0 = fixed interface method
С
                                              1 = free interface method
C
```

```
3 = residual-flexible free interface method
С
                                                  4 = user defined method
С
                                             - flags mixed interface method derived from
                                    cmsMixF
C
                                                  1 = free interface method
С
С
                                                  3 = Residual-flexible free interface method
                                                  0 = default
C
                                     kStress - key if modal element results are on file
С
                           ptrNARl,ptrNARh - 64 bit pointer to nodal averaged result records
С
                           ptrIRFl,ptrIRFh - 64 bit pointer to inertia relief modes
C
C
                           ptrNORl,ptrNORh - 64 bit pointer to normal modes
                           ptrCSTl,ptrCSTh - 64 bit pointer to constraint modes
С
                           ptrRSDl,ptrRSDh - 64 bit pointer to residual modes ptrELDl,ptrELDh - 64 bit pointer to element records
С
C
                                             - position not used
                                      0
C
С
             i
                      1
                           neqn
                                      SOLVER-to-ANSYS mapping vector (lsOLVtoANS(i), i=1,neqn)
                                      Note: When using the residual-flexible free interface method
С
С
                                            the modes are written in ANSYS internal equation ordering
                                            and the SOLVER-to-ANSYS mapping is not available. For
С
                                            other methods the modes are written in solver equation
C
                                            ordering and the SOLVER-to-ANSYS mapping is available.
С
                      1
                           lenbac
                                      Nodal equivalence table. This table equates the number used
C
                                      for storage to the actual node number.
C
                                      (Back(i), i=1, lenbac)
С
   NOR
                                      Normal Modes
C
            dp
                  nnorm
                           nean
С
                                      Inertia Relief Modes
С
   TRF
            dp
                  nirfm
                           negn
C
                                      Constraint Modes
С
   CST
                  ncstm
С
                                      Residual modes
С
   RSD
                 nrsdm
            dp
                           negn
c modal element results if kStress = 1:
С
            int
                    1
                           15
                                      nelm,
                                              maskl, nItems, ptrELM, ptrERS,
С
                                    ptrCER, ptrCERh, ptrESL, ptrESLh,
                                                                           nRF,
                                                                                     (10)
                                     ptrFR, ptrRFh, PrecKey,
                                                                 maskh,
C
                                     each item in header is described below:
С
                                       nelm - number of elements
C
                                    maskl,h - output mask (OUTRES)
C
С
                                     nItems - number of element records (7, VOL
                                              not included)
C
                                     ptrELM - pointer to element equivalence table
C
С
                                     ptrERS - pointer to element record sizes
С
                                   ptrCER,h - pointer to constant element records
                                   ptrESL,h - pointer to element index
C
                                        nRF - number of reaction forces
С
                                    ptrRF,h - pointer to reaction forces
С
                                    PrecKey - 0, double precision 1, single
C
                        above pointers are relative to ptrELD
С
C
            int
                         2*nItems
                                    Total size of each element record (LONGINT)
                                     Element equivalence table
С
   ELM
            int
                     1
                           nelm
                                      This table equates the order number used to
C
С
                                      the actual element number
С
   ERS
            int
                   nItems
                           nelm
                                     Sizes of the nItem element results sets for
С
                                      each element
   CER
                     1
                           5
                                   ptrVOL, ptrEPT, ptrEUL,
                                                                   0,
                                                                           0
С
            int
                        above pointers are relative to ptrCER
С
C
      constant element records (do not vary by mode):
    VOL
                     1 nelm*1
                                      Element volume
C
            dр
С
    EPT
            dр
                     1 nelm*size
                                      Element structural nodal temperatures
    EUL
            dр
                     1 nelm*size
                                     Element Euler angles
```

```
ESL
                                  ptrENS, ptrEEL, ptrEMS, ptrENF, ptrENG,
            int
                    1
                          10
C
                                  ptrENSh,ptrEELh,ptrEMSh,ptrENFh,ptrENGh
C
                       above pointers are relative to ptrESL
С
      non-constant element records (do vary by mode). Modes order is:
C
      --- nmode = [NOR CST (BCLV)]
                                         for fix interface method
С
      --- nmode = [NOR (IRF) CST (BCLV)] for free interface method
С
      --- nmode = [NOR RSD]
                                         for RFFB method
C
      --- () = if any
      --- BCLV = static correction vectors stored in .BCLV file
С
С
С
    ENS
            dp
                 nelm nmode*size
                                    Element nodal component stresses
                 nelm nmode*size
                                    Element nodal component elastic strains
C
    EEL
            dр
                 nelm nmode*size
                                    Element summable miscellaneous data
С
            dр
                                    Element nodal forces
С
    ENF
            dр
                 nelm nmode*size
    ENG
                 nelm nmode*3
                                    Element energies
С
            dp
С
      see fdresu.inc for more information on the element results
  NAR
            int
                    1
                           7
                                     nnod, ptrNCTl, ptrNCTh, ptrNSTl, ptrNSTh,
C
                                     ptrNELl, ptrNELh
C
                                    each item in header is described below:
C
                                     nnod - number of nodes
С
                                ptrNCTl,h - pointer to Node Contributions
С
                                ptrNSTl,h - pointer to NAR stress record
C
                                ptrNELl,h - pointer to NAR elastic strain record
С
                       above pointers are relative to ptrNAR
C
      NAR records (do vary by mode):
С
                nnod 2
C
    NCT
            i
                                    Node Contributions
    NST
                 nnod nmode*6
                                    Nodal component stresses
С
            dp
С
    NEL
            dp
                 nnod nmode*7
                                    Nodal component elastic strains
      see fdresu.inc for more information on the NAR records
```

1.8.3. TCMS File Format

```
*comdeck,fdtcms
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
      ******* description of CMS (component mode synthesis) transformation file ********
      character*8 TCMSNM
     parameter
                (TCMSNM='tcms
                     tcmsfpL, tcmsfp
     LONGINT
                     tcmsbk, tcmsut
     integer
      common /fdtcms/ tcmsfpL, tcmsbk, tcmsut
      equivalence (tcmsfp,tcmsfpL)
C
   ******* common variable descriptions *******
co tcmsfp
                file position on file tcms
   tcmsbk
                block number for file tcms
CO
    tcmsut
                file unit for file tcms
    See fddesc for documentation of how binary files are stored.
С
C
С
      ****** file format *******
        recid tells the identifier for this record. Not all records will have
C
             identifiers -- they are only indicated for those records whose
C
```

```
C
              record pointers are stored in the second file header.
         type tells what kind of information is stored in this record:
C
              i - integer
С
С
              dp - double precision
              cmp - complex
C
         nrec tells how many records of this description are found here
С
         lrec tells how long the records are (how many items are stored)
c recid
           type
                   nrec
                            lrec
                                     contents
                                     standard ANSYS file header (see binhed8 for
                            100
C
                    1
                                     details of header contents)
С
                             40
                                      .TCMS FILE HEADER
С
                    1
С
С
                                    fun48, nNodes,
                                                        neqn,
                                                              numdof, nirfm,
                                                         0,
                                                                    0,
                                    nnorm, ncstm,
C
                                  ptrNOR1, ptrCST1, ptrIRF1,
                                                                     0,
С
                                        0,
                                                0,
                                                          0,
                                                                     0,
                                                                              0,
C
                                  ptrNORh, ptrCSTh, ptrIRFh,
                                                                     0,
                                                                              0,
С
                                                  0,
                                        0,
                                                           0,
                                                                     Ο,
С
                                        0,
                                                  0,
                                                                    0,
                                                           0,
                                                                              0,
C
                                        0,
                                                  0,
С
                                                           0,
                                                  0,
С
                                        0,
                                                           0,
                                                                     0,
                                    each item in header is described below:
С
                                     fun48 - unit number
С
                                    nNodes - number of nodes in file
C
                                      neqn - number of equations (nNodes*numdof)
C
C
                                    numdof - number of dofs per node
                                     nirfm - number of inertia relief modes nnorm - number of normal modes
С
С
                                     ncstm - number of constraint modes
С
С
                           ptrIRFl,ptrIRFh - 64 bit pointer to inertia relief modes
С
                           ptrNORl,ptrNORh - 64 bit pointer to normal modes
                           ptrCSTl,ptrCSTh - 64 bit pointer to constraint modes
С
С
                                             - position not used
             i
                      1 nNodes
                                     Nodal equivalence table. This table equates
С
                                     the number used for storage to the actual
С
С
                                     node number
    NOR
                 nnorm
                                     Normal Modes
                           negn
                 nirfm
                                     Inertia Relief Modes
C
                           negn
    CST
                                     Constraint Modes
            ďρ
                 ncstm
                           nean
```

1.9. Description of the Full Stiffness-Mass File

This section explains the contents of the full file (jobname.full).

1.9.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 4.

1.9.2. FULL File Format

```
*comdeck,fdfull
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
      ******* description of full stiffness-mass file ********
      character*8 FULLNM
      parameter (FULLNM='full
С
      *** NOTE: if this variable is changed in the future it should be
                updated in spdefines.h also for symbolic assembly (jrb)
С
      integer
                      FULLHDLEN
                     (FULLHDLEN=160)
      parameter
      LONGINT
                      fullfpL, fullfp
      integer
                      fullbk, fullut, wrLdstep, wrSbstep, wrEqiter,
                      wr0ption
     common /fdfull/ fullfpL, fullbk, fullut,
                       wrLdstep,wrSbstep,wrEqiter,wrOption
      equivalence (fullfp,fullfpL)
    ****** common variable descriptions ********
C
  fullfpL
                 file position on file full
CO
   fullbk
                 block number for file full
   fullut.
                 file unit for file full
CO
      ****** file format (except for extopt=3,4) ********
С
    See fddesc for documentation of how binary files are stored.
С
      ****** file format *******
C
         recid tells the identifier for this record. Not all records will have
С
              identifiers -- they are only indicated for those records whose
C
С
              record pointers are stored in the second file header.
С
         type tells what kind of information is stored in this record:
              i - integer
С
              dp - double precision
C
              cmp - complex
C
         nrec tells how many records of this description are found here
C
         lrec tells how long the records are (how many items are stored)
c recid
                            lrec
                                     contents
           type
                   nrec
                    1
                            100
                                     standard ANSYS file header (see binhed8 for
С
С
                                     details of header contents)
            i
                    1
                            160
                                     .FULL FILE HEADER
C
С
                                    fun04.
                                                       nmrow,
                                                                 nmatrx,
                                                                                kan.
C
                                               negn,
                                             lenbac, numdof, ntermKl, ntermKh,
nmrow, ntermK, keyuns, extopt,
С
                                    wfmax,
С
                                    lumpm,
                                    keyse, sclstf, nxrows, ptrSTFl, ptrSTFh,
                                                                                        (20)
C
C
                                  ncefull, ntermMh, ptrENDl, ptrENDh, ptrIRHSl,
                                 ptrIRHSh, ptrMASl, ptrMASh, ptrDMPl, ptrDMPh,
С
                                                                                        (30)
                                  ptrCEl, ptrCEh, nNodes, ntermMl, ntermDl,
ptrDOFl, ptrDOFh, ptrRHSl, ptrRHSh, ntermDh,
С
С
                                  ngMaxNZ, ptrNGPHl, ptrNGPHh, minKdiag, maxKdiag,
C
С
                                 minMdiag, maxMdiag, minDdiag, maxDdiag, ngTerml,
                                                                                        (50)
С
                                  ngTermh, ngTermCl, ngTermCh,ptrDIAGKl,ptrDIAGKh,
С
                                ptrDIAGMl,ptrDIAGMh,ptrDIAGCl,ptrDIAGCh, ptrSCLKl,
                                                                                        (60)
                                ptrSCLKh, Glbneqn, distKey, ngTermFl, ngTermFh,
GlbnNodes, GlbnVars, GlbfAcCE, lcAcLen, GlbfCE,
С
C
                                                                                        (70)
```

```
С
                                    ptrGmtl, ptrGmth,nceGprime,numA12A11,GnVirtBCs,
                                    ntermGl, ntermGh,ptrDensel,ptrDenseh, nVirtBCs,
С
                                                                                           (80)
                                 ptrVrtBCl,ptrVrtBCh, ptrMRKl, ptrMRKh, ptrKclxl,
С
                                   ptrKclxh, ntermKCl, ntermKCh,minKCdiag,maxKCdiag,
С
С
                                      ngChg, ptrBCl, ptrBCh, ptrPHYSl, ptrPHYSh,
                                    predKey,fullDistF, ptrGVBCl, ptrGVBCh,
                                                                                nzRow,
C
                                                                                          (100)
                              localNonlKey, nMastDOF, ptrMDFl, ptrMDFh, GlbnMast,
    ptrGMDFl, ptrGMDFh, cmsMeth, cmsMixF, hrmopt,
С
С
                                                                                          (110)
                                    ActFlag, InActNKey, IANSndCnt, IANRcvCnt, Glblenbac,
C
C
                                    ptrActl, ptrActh,ext_nNods, ext_neqn,ext_nmrow,
                                                                                          (120)
                                MtxEqnFlag,
                                                    0,
                                                               0,
                                                                          0,
                                                                                     0,
C
С
                                          0,
                                                     0,
                                                                Ο,
                                                                           0,
                                                                                      0,
                                                                                          (130)
C
                                          0,
                                                     0,
                                                               0,
                                                                          0,
                                                                                     0,
                                          0,
                                                     0,
                                                               0,
                                                                          0,
                                                                                     0,
                                                                                          (140)
C
C
                                          0,
                                                     0,
                                                                Ο,
                                                                           0,
                                                                                      0,
С
                                          0,
                                                     0,
                                                               0,
                                                                          0,
                                                                                     0,
                                                                                          (150)
                                          0,
                                                     0,
                                                                0,
                                                                           0,
                                                                                     0,
С
                                                                0,
                                                                           0,
                                                                                          (160)
С
                                          0,
                                                     0,
                                                                                      0
                                      each item in header is described below:
C
                                       fun04 - negative of the unit number (-4)
С
                                                 NOTE: if fun04 is > 0 it means that
C
                                                       frontal assembly was used (which
C
                                                       is longer documented here)
С
С
                                              - number of equations on file
C
                                       nmrow - number of active DOF (neqn-BC)
                                       nmatrx - number of matrices on file
С
                                              - analysis type
C
                                       kan
                                       wfmax - maximum row size
С
                                       lenbac - number of nodes in ANSYS space (this can be different from nNodes wh
C
C
                                       numdof - number of dofs per node
                                       ntermKl,ntermKh - number of terms in Stiffness
C
С
                                                          matrix
                                       lumpm - lumped mass key
C
С
                                                 = 0 - default matrix type
                                                = 1 - lumped
С
                                       ntermK - pre-8.1 this is the number of terms
C
                                                 in Stiffness matrix (otherwise this
С
                                                value must be 0 and ntermKl,ntermKh
С
                                                must be used)
С
                                       keyuns - unsymmetric key
C
C
                                                 = 0 - no unsymmetric matrices on
С
С
                                                 = 1 - there is at least one
                                                        unsymmetric matrix on file
C
С
                                       extopt - mode extraction method
С
                                                 = 0 - reduced
                                                 = 1 - lumped
C
                                                 = 3 - unsymmetric Lanczos
С
                                                 = 4 - damped Lanczos
С
                                                 = 6 - block Lanczos
C
                                                 = 7 - QRdamped
C
                                                 = 8 - SuperNode
C
С
                                                = 9 - PCG Lanczos
                                       keyse - superelement key; set if at least
С
C
                                               one superelement
                                       sclstf - maximum absolute stiffness matrix term
С
С
                                       nxrows - the maximum rank for this solution
                                       ptrSTFl,h - pointer to Stiffness matrix
C
                                                  number of CE+CP equationslow part of 64 bit end of file ptr
С
С
                                       ptrENDl
                                                  - high part of 64 bit end of file ptr
C
                                       ptrENDh
                                       ptrIRHSl,h - pointer to imaginary RHS (F)
C
                                       {\tt ptrMASl,h} \  \, {\tt -pointer} \  \, {\tt to} \  \, {\tt Mass} \  \, {\tt matrix}
C
С
                                       ptrDMPl,h - pointer to Damping matrix
С
                                       ptrCEl,h - pointer to Gt and g matrices
                                                   - number of nodes considered by assembly (nNodes can be different
C
                                       ntermMl,h - number of terms in Mass matrix
С
                                       ntermDl,h - number of terms in Damping matrix
```

С	ptrDOF1,h - pointer to DOF info
С	ptrRHSl,h - pointer to RHS (F)
С	ngMaxNZ - maximum number of nodes per nodal
C	block in nodal graph structure
C	ptrNGPH1,h - pointer to vectors needed for nodal graph structure
C	minKdiag - minimum absolute stiffness matrix
c	diagonal term
C	maxKdiag - maximum absolute stiffness matrix
C	diagonal term
C	minMdiag - minimum absolute mass matrix
С	diagonal term
C	maxMdiag - maximum absolute mass matrix
С	diagonal term
C	minDdiag - minimum absolute damping matrix
C	diagonal term
С	maxDdiag - maximum absolute damping matrix
С	diagonal term
С	ngTerml,h - total number of nonzeroes in nodal graph
C	(expanded graph based value, no BC applied)
C C	ngTermCl,h - total number of nonzeroes in nodal graph (compressed graph based value)
C	ptrDIAGKl,h - pointer to stiffness matrix DIAGONAL vector
C	(NOTE: this is a copy of the diagonal
c	values stored in the full matrix)
C	ptrDIAGMl,h - pointer to mass matrix DIAGONAL vector
C	(NOTE: this is a copy of the diagonal
С	values stored in the full matrix)
С	ptrDIAGCl,h - pointer to damping matrix DIAGONAL vector
С	(NOTE: this is a copy of the diagonal
С	values stored in the full matrix)
С	ptrSCLK1,h - pointer to stiffness matrix diagonal scaling
C	vector (may contain all 1.0's when not scaling)
C	Glbneqn - global number of active DOF (this will match nmrow
C	unless we are writing distributed "local" FULL files
C	in Distributed ANSYS
C	distKey - key denoting whether the FULL file is a single,
C	global FULL file (0) or multiple, local FULL file (1) ngTermFl,h - total number of nonzeroes in nodal graph
C	as passed to the solver (after BC applied)
c	GlbnNodes - global number of nodes considered by assembly
C	GlbnVars - global number of equations (will match neqn
C	unless we are writing distributed "local" FULL files
C	in Distributed ANSYS
С	GlbfAcCE - total number of across CE
С	lcAcLen - number of acrossCE where depedents are in my domain
С	GlbfCE - total number of all the CE
C	ptrGmtl,h - pointer to G prime matrix for local nonlinearity
С	nceGprime - number of CE (or equations) in G prime local nonlinearit
С	numA12A11 - number of equations in local nonlinear matrix: excluding
C	equations from G prime
C	GnVirtBCs - global number of virtual constraints
C	ntermGl,ntermGh - total number of terms in total local nonlinear
C	matrix including A12,A11 and G: total sum
C	ptrDensel,ptrDenseh - dense matrix information in local nonlinear matrix
C	matrix nVirtBCs - number of virtual constraints
C	ptrVrtBCl,ptrVrtBCh - pointer to the virtual constraint DOF data
C	ptrMRKl,h - pointer to the DOF marker array
C	0 - position not used
C	ptrKclxl,h - pointer to K complex (the 4th matrix) matrix
C	full case is: K, M, C, Kcplx: existing at same time
С	ntermKCl,h - number of terms in Complex Stiffness matrix
C	minKCdiag - minimum absolute complex stiffness matrix
C	diagonal term
С	maxKCdiag - maximum absolute complex stiffness matrix
С	diagonal term
С	ngChg - key denoting whether or not the nodal graph written
С	to this FULL file differs from the previous FULL file
С	ptrBCl,h - pointer to boundary condition data
C	ptrPHYSl,h- pointer to the physics marker array

```
С
                                     predKey - nonlinear analysis predictor key
                                     fullDistF - indicate the global profiling of distributed full file
С
                                     ptrGVBCl,h- pointer to the global virtual constraint DOF data
C
                                               - number of non-zero rows of the matrics for the full file under
С
                                     nzRow
С
                                                 global full file, -1 means not applicable
                                     localNonlKey - local nonlinearity speedup key
C
                                              - number of master DOF for substructuring
С
                                     ptrMDFl,h - pointer to the master DOF data for substructuring
С
                                     GlbnMast - global number of master DOF for substructuring
C
C
                                     ptrGMDFl,h- pointer to the global master DOF data for substructuring
                                     cmsMeth - CMS method key
С
С
                                                 = 0 - fixed interface method
                                                 = 1 - free interface method
С
                                                 = 3 - residual-flexible free interface method
С
                                                 = 4 - user defined method
С
С
                                     cmsMixF
                                               - flags mixed interface method derived from
                                                 = 1 - free interface method
С
                                                 = 3 - Residual-flexible free interface method
= 0 - default
С
С
                                                - harmonic analysis type
С
                                     hrmopt
                                                 = 0 Not a harmonic analysis
C
                                                 > 0 Harmonic analysis type
С
С
                                     ActFlag
                                                - active element logic flag for the two-step method (only assemble
                                                - equivalent to actSymbFlag defined in soptcm.inc
С
                                                - currently, it is used for AM and target element trimming
C
С
                                                  = 2 - mark active nodes in casiInitialize before assembly due to
С
                                                  = 1 - mark active nodes in casiInitialize before assembly due to
С
                                                  = 0 - default
                                     InActNKey - sum of number of inactive nodes across all domain.
С
                                                - it is only useful when ActFlag = 2
С
                                                  = 0 - number of active nodes (nNodes) is equal to original ANSYS
С
                                                      - there is no need to do AllToAll communication for the inac
C
C
                                                  > 0 - number of active nodes (nNodes) is smaller than original A
                                                       - if distributed full file is used, AllToAll communication a
С
С
                                                      - back to ANSYS dof space
                                     IANSndCnt - total number of inactive nodes that need to be sent to other dom
C
С
                                                - it is only usefull when InActNKey > 0 and distributed full file
С
                                                  = 0 - no inactive nodes information in current domain that need
                                                  > 0 - length of InActNodesSnd array
С
                                     IANRcvCnt - total number of inactive nodes that need to be received to other
С
                                                - it is only usefull when InActNKey > 0 and distributed full file
С
                                                  = 0 - no inactive nodes information in current domain that need
С
                                                  > 0 - length of InActNodesRcv array
C
С
                                     Glblenbac - Global number of nodes in ANSYS space
С
                                     ptrActl,h - pointer to act nodes to ANSYS nodes mapping array and other supp
С
                                     ext_nNods - nNodes + extra number of nodes needed by local equation based ma
                                     ext_neqn - neqn on file + extra number of equations needed by local equatio
C
С
                                     ext_nmrow - ext_neqn - nBC - number of BC in extra number of equations neede
С
                                     EqnMtxFlag - flag indicate whether we are using equation based matrix in full
                                                  = 0 - element based full file (default)
C
                                                  = 1 - equation based full file
С
С
C
                           numdof
                                     Degrees of freedom per node
С
                                     DOF reference numbers are:
C
C
          UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8
          AZ = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
C
          CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
C
С
          EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
                    1
                                     Nodal equivalence table. This table equates
C
                           lenbac
С
                                     the number used for storage to the actual
С
                                     node number
                    1
                        Glblenbac
                                     Global nodal equivalence table. This record
С
                                     EXISTS ONLY in the full file of the master
С
                                     domain and ONLY for distributed .full files
C
\ensuremath{\mathtt{c}} Stiffness Matrix. The next two records are repeated as a group neqn times.
c The pair of records will repeat GlbnVars times when model has across CEs.
c And row indices are global in the case.
```

C C C	STF	i	1	varies	Matrix row indices. The last item corresponds to the diagonal. The length of this record will vary (actual length is returned from routine BINRD8)		
С		dp/cmp	1	varies	Matrix terms		
C C					If keyuns=0, this record will contain the terms before the diagonal.		
C C C	Load Ve	ector			If keyuns=1, this record will contain the entire row.		
С	RHS	dp/cmp	1	negn	Load vector terms.		
С		ary part of		_			
	IRHS	dp	1		Imaginary load vector terms.		
c c c	CHAI	αр	Ţ	neqn	This record EXISTS ONLY if its pointer in the header is not zero. Record length will be GlbnVars for model with across CE.		
С	Stiffne	ess matrix	diag	onal vector			
C C	DIAGK	dp/cmp	1	neqn	diagonal vector data for stiffness matrix. Its length will be GlbnVars for model with across CE.		
С	Stiffne	ess matrix	diag	onal scaling	vector		
C C	SCLK	dp/cmp	1	ext_neqn	diagonal scaling vector for stiffness matrix. Record length will be GlbnVars for across CE model.		
C	DOF mar	rker array					
0 0 0 0 0	MRK	i	1	neqn	marker array flagging various types of DOF (1=U_EQN, 2=P_EQN, 3=E_EQN, 4=A_EQN). Positive values mean the DOF belongs to a user-defined node, negative values mean the DOF belongs to an internal node. NOTE: if this array does not exist then it is assumed that all DOFs are U_EQNs for user-defined nodes. Record length will be GlbnVars for models with across CE.		
С	PHYSICS	3 marker ar	rray				
0 0 0 0 0	PHYS	i	1	neqn	marker array flagging the various types of physics (1=STRUCTURAL_EQN, 2=THERMAL_EQN, 3=ELECTRICAL_EQN, 4=MAGNETIC_EQN, 5=FLUID_EQN, 6=DIFFUSION_EQN) NOTE: if this array does not exist then it is assumed that all DOFs are STRUCTURAL_EQNs. Record length will be GlbnVars for models with across CE.		
c DOF information							
С	DOF	i	1	ext_nNods	Nodal extent vector. Number of DOFs at each node		
с с		i	1	GlbnNodes	Global nodal extent vector giving numbers of DOFs at each global nodes. This record EXISTS ONLY for models using DMP with across CE.		
с с		i	1	GlbnNodes	A vector mapping global node number to local node number1 in the vector means the node is not in this domain. This record EXISTS ONLY for model with across CE.		
С		i	1	ext_neqn	DOF vector. If negative, this DOF is constrained.		
C		i	1	GlbnVars	A vector of global DOF reference numbers. this record EXISTS ONLY for models using DMP with across CE.		

```
C
                        ext_nNods
                                     A vector mapping local node number to global node number.
                                     This record EXISTS ONLY for distributed .full files
С
c act nodes to ANSYS nodes mapping array
            i
                    1
                         ext_nNods A vector mapping active nodes used in assembly to ANSYS nodes.
C
                                    A vector mapping global active nodes to global ANSYS nodes
С
                    1
                         GlbnNodes
                                     This record EXISTS ONLY for models using DMP with across CE.
C
                    1
                           numCPII
                                     A vector (InActNumNodesSnd) giving the number of inactive nodes
C
            i
C
                                     that needs to be sent to each domain
C
                                     This record EXISTS ONLY for distributed .full files and InActNKey > 0
                                     A vector (InActNumNodesRcv) giving the number of inactive nodes
C
                    1
                           numCPU
С
                                     that needs to be received from each domain
                                     This record EXISTS ONLY for distributed .full files and InActNKey > 0
C
С
                    1
                         IANSndCnt
                                     A vector (InActNodesSnd) giving local internal node number that needs to be
                                     sent to other domains.
C
                                     This record EXISTS ONLY when IANSndCnt > 0
C
                         IANRcvCnt
                                     A vector (InActNodesRcv) giving local internal node number that needs to be
C
                    1
                                     received from other domains.
C
                                     This record EXISTS ONLY when IANRcvdCnt > 0
C
c BC information
С
  BC
                    1
                             neqn
                                     DOFs with imposed values
                    1
                            varies
         dp/cmp
                                     Imposed values
C
c Mass Matrix.
     if lumpm = 0:
C
       The next two records are repeated as a group neqn times.
       It will be in global form the same way as stiffness matrix if model has across CE.
C
С
  MAS
                    1
                            varies
                                     Matrix row indices. The last item
                                     corresponds to the diagonal. The
C
                                     length of this record will vary (actual
С
                                     length is returned from routine BINRD8)
С
                            varies
                                     Matrix terms
C
           dp
                    1
     if lumpm = 1:
С
                    1
                              neqn
                                     Matrix diagonals.
C
           dp
                                     Record length will be GlbnVars for across CE model.
c Mass matrix diagonal vector
                    1
                                     diagonal vector data for mass matrix.
С
   DIAGM
           ďρ
                              neqn
                                     Record length will be GlbnVars for across CE model.
C
c Damping Matrix. The next two records are repeated as a group neqn times.
c For model with across CE, it will be in global form the same way as stiffness matrix.
С
   DMP
                    1
                            varies
                                     Matrix row indices. The last item
                                     corresponds to the diagonal. The
C
С
                                     length of this record will vary (actual
С
                                     length is returned from routine BINRD8)
C
           dр
                    1
                            varies
                                     Matrix terms
c Damping matrix diagonal vector
                                     diagonal vector data for damping matrix.
С
  DIAGC
           dp
                    1
                              negn
                                     Record length will be GlbnVars for across CE model.
C
c K complex Matrix. The next two records are repeated as a group neqn times.
c For model with across CE, it will be in global form the same way as stiffness matrix.
```

C C C	KC	i	1	varies	Matrix row indices. The last item corresponds to the diagonal. The length of this record will vary (actual length is returned from routine BINRD8)			
С		dp	1	varies	Matrix terms			
c Nodal graph vectors								
С	NGPH	i	1	nNodes	number of nonzeroes for each node.			
	c Record length will be GlbnNodes for across CE model c Repeat for each node							
С	2	i	1	varies	Index vector. Node number in the vector is global when			
С	Q		£11 .	0	model has across CE			
С	G matriz	x 11 nce	itull >	0.				
C	CE	i	1	ncefull	List of dependent DOFs of local CEs. It EXISTS ONLY if ncefull>0. The dependent DOF is local			
С С		i	1	lcAcCE	List of dependent DOFs of local across CEs. This record EXISTS ONLY if lcAcCE>0. The dependent DOF is local			
C C		i	1	GlbfAcCE	List of dependent DOFs of all across CEs. This record EXISTS ONLY if GlbfAcCE>0. And it is ONLY in the full file of master domain			
CCC		i	1	GlbfCE	List of dependent DOFs of all CEs. This record EXISTS ONLY if GlbAcCE>0. And it is ONLY in the full file of the master domain			
C		dр	1	ncefull	g vector (constant terms) of local CEs. This record EXISTS ONLY if ncefull>0			
CC		dp	1	ncefull	imaginary g vector (constant terms) of local CEs. This vector only exists for FULL harmonic analyses (kan=3). This record EXISTS ONLY if ncefull>0			
C		dp	1	ncefull	g vector (constant terms) of local CEs for nonlinear analysis predictor logic. This record EXISTS ONLY if ncefull>0 & predKey=1.			
		dn	1	GlbfAcCE				
CC		dp	1	GIDIACCE	g vector (constant terms) of across CEs. This record EXISTS ONLY if GlbfAcCE>0 in the full file of the master domain			
0 0 0		dp	1	GlbfAcCE	imaginary g vector (constant terms) of across CEs. This vector only exists for FULL harmonic analyses (kan=3). This record EXISTS ONLY if GlbfAcCE>0 in the full file of the master domain			
CCC		dp	1	GlbfAcCE	g vector (constant terms) of across CEs for nonlinear analysis predictor logic. This record EXISTS ONLY if GlbfAcCE>0 & predKey=1 in the master full file.			
C	Follow:	ing loca i	.1 CE da	ata records 4	EXIST ONLY in the full file of the domain with local CEs: Header for local CEs; 1=nRows, 2=nRows, 3=1, 4=0			
С		i	1	nRows	Vector of 1's			
С		i	1	nRows	Number of non-zero terms in each row for one local CE			
С	Repeat	t for ea	ch row					
С		i	1	varies	Column indices in local equation numbers			
С		dp	1	varies	Column values			

```
Following across CE data records EXIST ONLY if GlbfAcCE>0 in the full file of master domain:
            i
                    1
                                 4
                                     Header for across CEs; 1=nRows, 2=nRows, 3=1, 4=0
C
            i
                    1
                                     Vector of 1's
C
                             nRows
С
            i
                    1
                             nRows
                                     Number of non-zero terms in each row for an across CE
    Repeat for each row:
C
                            varies
                                     Column indices in global equation numbers
                                     Column values
C
           dр
                    1
                            varies
    Following CE data records EXIST ONLY if n>0 in the full file, where n == ncefull with
C
    SMP and n == GlbfCE with DMP. NOTE: for DMP these records only exist in the .full file
С
    of master domain
C
                                    List of dependent DOFs of all CEs. This record EXISTS ONLY
C
                                     if n>0. And it is ONLY in the full file of the master
C
                                     domain
С
    NOTE: this matrix includes boundary d.o.f. which touch the CEs (for cnvfor.F)
C
C
                    1
                                     Header for across CEs; 1=nRows, 2=nRows, 3=1, 4=0
            i
                    1
                             nRows
                                     Vector of 1's
C
                    1
                                     Number of non-zero terms in each row for an across CE
С
            i
                             nRows
    Repeat for each row:
            i
                    1
                                     Column indices in global equation numbers
                            varies
C
                    1
                            varies
                                     Column values
           ďρ
c Virtual constraint vector
  VBC
                    1
                          nVirtBCs
                                     marker array (bit 1 set -> constrained DOF for residual vector)
С
С
                                                   (bit 2 set -> constrained DOF for enforced motion)
                                                   (bit 3 set -> eliminated DOF for substructure master DOF)
С
            i
                    1
                         nVirtBCs
                                     virtual constraint DOFs
C
С
   GVBC
                          GnVirtBCs
                                    marker array (bit 1 set -> constrained DOF for residual vector)
C
                                                   (bit 2 set -> constrained DOF for enforced motion)
                                                   (bit 3 set -> eliminated DOF for substructure master DOF)
C
                                     this record EXISTS ONLY for models using DMP with across CE
С
            i
                    1
                          GnVirtBCs
                                    virtual constraint DOFs
C
                                     this record EXISTS ONLY for models using DMP with across CE
C
c Substructuring master DOF vector
                                     list of master DOFs for substructuring
C
  MDF
            i
                    1
                         nMast.DOF
С
   GMDF
                    1
                         GlbnMast
                                     list of master DOFs for substructuring
                                     this record EXISTS ONLY for models using DMP with across CE
C
  Meaning of K11, K12, and G matrices:
C
      Given
C
          [K]\{x\} = \{F\}
С
      subject to the constraints
С
C
            \{x1\} = [G]\{x2\} + \{g\}
      where {x1} are the dependent DOFs, {x2} the independent DOFs
C
     This results in
С
```

```
[K^*]\{x2\} = \{F^*\}
C
С
      where
          [K^*] = [G]'[K11][G] + [G]'[K12] + [K21][G] + [K22]
C
          {F*} = [G]'{f1} + {f2} - [G]'[K11]{g} - [K21]{g}
C
c complex version of \{F^*\} decomposed into, we assume G' is always real
c and g could be complex denoted as g' == (g,gx):
          G' K11' g' = G' (K11,M11)*(g,gx)
                     = G' [K11*g - M11*gx, M11*g + K11*gx]
C
          K21' *g' = (K21, M21)*(g,gx)
C
С
                      = (K21*g- M21*gx, K21*gx + M21*g)
```

1.10. Description of the Substructure Displacement File

This section explains the contents of the substructure displacement file (jobname.dsub).

1.10.1. Standard Mechanical APDL File Header

See The Standard Header for Mechanical APDL Binary Files (p. 4) for a description of this set. File number (Item 1) is 13.

1.10.2. DSUB File Format

```
*comdeck,fddsub
c *** Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
     ******* description of substructure displacement file ****
     character*8 DSUBNM
     parameter (DSUBNM='dsub
     LONGINT
                   dsubfpL, dsubfp
                  dsubbk, dsubut
     integer
     common /fddsub/ dsubfpL, dsubbk, dsubut
     equivalence (dsubfp,dsubfpL)
     ****************
С
                          CAUTION
С
        Please update proc getDSUBInfo in SEManagement.eui
С
С
        if the file format changes or GUI read of the DSUB
       file will fail.
C
     *****************
С
   ****** common variable descriptions ********
C
  dsubfpL
              file position on file dsub
             block number for file dsub
co dsubbk
co dsubut
              file unit for file dsub
c open: slvstr
         setdis,eostrt,eofini,ranbasPCG,ranbwvPCG,slvstr,slvend,supsrl,supidx,supscl
  write:
  close: slvend
c read: matstr as .usub
   See fddesc for documentation of how binary files are stored.
C
C
     ****** file format *******
C
```

```
recid tells the identifier for this record. Not all records will have
С
              identifiers -- they are only indicated for those records whose
С
              record pointers are stored in the second file header.
C
С
         type tells what kind of information is stored in this record:
              i - integer
C
              dp - double precision
cmp - complex
C
С
C
         nrec tells how many records of this description are found here
C
         lrec tells how long the records are (how many items are stored)
c recid
                                     contents
           type
                   nrec
                           lrec
                                     standard ANSYS file header (see binhed8 for
С
                    1
                            100
                                     details of header contents)
C
С
    1
            i
                    1
                             20
                                     .DSUB FILE HEADER
                                    fun13, fpeofS, fpeofL,
                                                               kcxp,
C
                                     knum, kCXFM, senres, cpxeng,
                                                                          0,
C
С
                                        0,
                                                Ο,
                                                         Ο,
                                                                  0,
                                                                          0,
                                        0,
                                                0,
                                                         0,
                                                                  0,
                                                                          0,
С
C
                              fun13 - file unit number
С
                      fpeofS, fpeofL - pointer to the eof
С
C
                              kcxp
                                      - = 1 if complex results
                                     - number of extracted modes if use pass modal
С
                                      - number of expanded modes if use pass modal
С
                              knum
                              senres - = 1: expand only displacements
C
С
                          (all type of analysis)
C
                                        = 2: expand real and then imaginary part
С
                                             of displacements (harmonic analysis)
С
                                        = 3: expand displacements, velocities,
                                             accelerations (transient analysis)
С
С
                              cpxeng - = 1: after use pass = harmonic or modal damp
С
                                             compute average, amplitude, peak energies
С
       *** these records are repeated each iteration ***
                                   fun13, kan, lenbac, numdof, kcmplx,
                             50
С
                    1
                                    itime, itter, ncumit, nitter, curdof(i),
С
                                                                                (10)
                                   (curdof(i),i=1,numdof)
С
С
                                   (curdof(i),i=1,numdof)
                                                                                 (20)
                                   (curdof(i), i=1, numdof)
С
С
                                   (curdof(i), i=1, numdof)
                                                                                 (30)
                                   (curdof(i),i=1,numdof)
С
С
                                   (curdof(i), i=1, numdof)
                                                                                 (40)
                                curdof(i),
С
                                            0, 0,
                                                                  0,
                                                                            0,
                                        0, extopt,qrDmpKy,Glblenbac, timint,
C
                                                                                (50)
С
                              extopt - mode extraction option
С
С
                              qrDmpKy - QR damped calculations key
    3
           ďρ
                    1
                             2.0
                                   time/freq,acel(1),acel(2),acel(3), frqenr,
C
                                                                0,
С
                                       frqud,
                                                   0,
                                                       0,
                                                                            0,
С
                                           0,
                                                    0,
                                                            0,
                                                                    0,
                                                                            0,
                                           0,
                                                                            0,
                                                    0,
                                                            0.
C
С
       *** the following records are repeated for each superelement ***
           i
                             20
                                   iel,
                                         nrow, nvect, ntrans, namel,
                 1
С
                                   name2, trok, lrok, name3, name4,
name5, name6, name7, name8, kCXFM,
С
С
                                                       0,
С
                                   kdamp,
                                               0,
                                                               0,
С
                              iel
                                     - (iel=0 signals end of superelements
С
С
                                       for this iteration)
С
                              nrow
                                     - number of dofs
                              nvect - number of load vectors
C
                              ntrans - number of transformations
С
                             name* - name1/2/3/4/5/6/7/8 are the
```

```
inexc4 representation of the
С
                                      32 character filename.
С
                                      name3/4/5/6/7/8 will be "0"
С
                                      for pre rev 5.2 files - cwa
С
                                    - flag if transformations can be applied
С
                             trok
                                    - flag if large deformation transformation
С
                             lrok
С
                                      can be applied
                             kCXFM - key if CXFM transformations are present on file
С
                             kdamp
                                   - key if ratios:
С
                                      rdamp(1),rdamp(2),dmprat,dmpst,spin,mscalm
С
                                      are present on file
С
С
С
          dр
                   1
                           125
                                  ntrans sets of transformations 15x5
С
С
          dр
                           250
                                  ntrans sets of transformations (CS) 50x5 - present if kCXFM = 1
   7
          LONG
                                 (lL(i),i=1,nrow) - global dofs
С
                    1
                          nrow
                                   (made LONGINT in version 14.0)
С
С
    8
          dр
                    1
                          nvect
                                  (scalf(i), i=1, nvect)
                                  rdamp(1),rdamp(2),dmprat,dmpstr,spin,
С
   9
          dр
                    1
                           10
                                                                           - present if kdamp = 1
                                  mscalm , 0,
С
                                                         0, 0, 0,
С
  10
           dр
                    1
                          nrow
                                  (disp(i),i=1,nrow)
  11
                                  (vel(i),i=1,nrow)
                                                                           - present if senres >= 2
С
          dр
                    1
                          nrow
  12
          dр
                    1
                          nrow
                                  (acel(i),i=1,nrow)
                                                                           - present if senres = 3
```

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Chapter 2: Accessing Binary Data Files

This chapter explains the routines you need to read, write, or modify a Mechanical APDL binary file. This collection of routines (called BINLIB) resides on your product-distribution media.

The following topics are discussed in this chapter:

- 2.1. Accessing Mechanical APDL Binary Files
- 2.2. Demonstration Routines
- 2.3. Retrieving Data from the Results File

2.1. Accessing Mechanical APDL Binary Files

The BINLIB library is in the dynamic link library \Program Files\ANSYS Inc\V211\ANSYS\custom\misc\<platform>\binlib.dll on Windows systems (where <platform> is a directory that uniquely identifies the hardware platform version) or the shared library /ansys_inc/v211/ansys/customize/misc/<platform>/libbin.so on Linux systems.

2.1.1. Access Routines for Results Files

Demonstration programs that use the BINLIB library for reading and writing Mechanical APDL results files are included on the installation media:

- ResRdDemo
- ResWrDemo
- · bintst

On Windows Systems:

The FORTRAN source for these programs is located in \Program Files\ANSYS Inc\V211\ANSYS\customize\user. The files are named ResRdDemo.F and ResWrDemo.F.

To link these demonstration programs, use the \Program Files\ANSYS Inc\V211\ANSYS\custom\misc\<platform>\rdrwrt.bat procedure file and specify the program that you want to build on the command line. Valid command line options are ResRdDemo, ResWrDemo, and user-prog. For example, to build the program to read a results file, type:

\Program Files\ANSYS Inc\V211\ANSYS\custom\misc\<platform>\rdrwrt ResRdDemo

Appropriate files are then copied from \Program Files\ANSYS Inc\V211\ANSYS\custom-ize\user to your working directory, compiled, and linked. The resulting executable is also placed in your current working directory.

Use the userprog command line option when writing your own customized program, naming the routine userprog.F. The resulting executable is named userprog.exe. When userprog is used, no files are copied from $\program\progr$

These files are loaded onto your system only if you performed a custom installation and chose to install the customization tools.

On Linux systems:

The FORTRAN source for these programs is located in /ansys_inc/v211/ansys/custom-ize/misc. The files are named ResRdDemo.F and ResWrDemo.F.

To link these demonstration programs, use the /ansys_inc/v211/ansys/custom-ize/misc/rdrwrt.link procedure file and specify the program that you want to build on the command line. Valid command line options are ResRdDemo, ResWrDemo, and userprog. For example, to build the program to read a results file, type:

/ansys_inc/v211/ansys/customize/misc/rdrwrt.link ResRdDemo

Appropriate files are then copied from /ansys_inc/v211/ansys/customize/misc to your working directory, compiled, and linked. The resulting executable is also placed in your current working directory. Procedure files are available in the /ansys_inc/v211/ansys/bin directory to run these programs, once linked. The procedure files are named ResRdDemo211 and ResWr-Demo211.

Use the userprog command line option when writing your own customized program, naming the routine userprog.F. The resulting executable is named userprog.e211. When userprog is used, no files are copied from /ansys_inc/v211/ansys/customize/misc to your working directory. The procedure file is named userprog211.

These files are loaded onto your system only if you performed a custom installation and chose to install the customization tools.

2.1.2. Characteristics of Mechanical APDL Binary Files

Before accessing Mechanical APDL binary files, you need to know certain file characteristics:

- 1. A Mechanical APDL binary file is a direct-access, unformatted file. You read or write a record by specifying (as a number) what location to read or write.
- 2. Before Mechanical APDL actually writes data to a file on a disk, it uses buffers to store data in memory until those buffers become full. A block number designates these buffers. Most access routines use this block number.
- 3. By default, Mechanical APDL files are external files. The standardized "external" format the files use enables you to transport them across different computer systems.
- 4. In addition to file names, Mechanical APDL uses file numbers to identify the files. File handles and other information are associated with the file numbers.

- 5. Some binary files contain data values that point to the start of certain data (for example, the start of the data steps index table record). Both Mechanical APDL and external binary files access routines use these pointers to locate data on the various binary files.
- 6. All data is written out as 32-bit integers. Double-precision data and pointers, therefore, take up two integer words. To create a 64-bit pointer from the two 32-bit integers, use the function largeIntGet.

2.1.3. Viewing Binary File Contents

To view the contents of certain Mechanical APDL binary files, you issue the command /AUX2 or choose menu path Utility Menu>File>List>Binary Files or Utility Menu>List>File>Binary Files. (You can do so only at the Begin level.) Mechanical APDL then enters its binary file dumping processor, AUX2, and dumps the binary file record by record.

In AUX2, you can use either the record number (**DUMP** command) or the record pointer (**PTR** command). If the file was written in parallel (-NP>1 on the command line), the **DUMP** command may not work as expected. In that case, only the **PTR** command may be used.

2.1.4. Abbreviations

The input and output for the routines discussed in this chapter are described with the following abbreviations:

• Type of variable is one of the following:

```
int - integer (4-byte)
long - integer (8-byte)
dp - double-precision
log - logical (true or false)
char - character
comp - double precision complex
```

• *Size* of variable is one of the following:

```
sc - scalar variable
ar(n) - array size n
func - functional return value
```

• *Intent* of variable is one of the following:

```
in - input onlyout - output onlyinout - both an input and an output variable
```

2.1.5. binini (Initializing Buffered Binary I/O Systems)

```
*deck,binini
subroutine binini (iott)

c *** primary function: initialize buffered binary i/o system

c --- This routine is intended to be used in standalone programs.

c --- This routine should not be linked into the ANSYS program.

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
    iott (int,sc,in) - output unit number for error output

c output arguments: none
```

2.1.6. Function sysiqr (Retrieving the Status of a File)

```
*deck,sysiqr
     function sysiqr (nunit,fname,lname_in,inqr_in)
c *** primary function: do a file system inquire (system dependent)
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
    variable (typ,siz,intent) description
   nunit (int,sc,in)
                              - fortran unit number (used only for inqr='0')
C
   fname (chr,sc,in)
                             - name of file
С
  lname_in (int,sc,in)
                              - length of file name (characters, max=50)
   inqr_in (chr,sc,in)
                              - character key for information requested
C
С
                                 = 'E' - return whether file exists
                                     sysiqr = 1 - file exists
С
                                           = 0 - file does not exist
C
                                            < 0 - error occured
С
                                 = '0' - return whether file is open
С
C
                                     sysigr = 1 - file is open
                                            = 0 - file is closed
С
                                            < 0 - error occured
C
                                 = 'N' - return unit number of file
С
С
                                     sysigr > 0 - unit number for file
                                            = 0 - file not assigned to a unit
С
                                            < 0 - error occured
С
C
 output arguments:
С
    sysigr (int,func,out) - the returned value of sysigr is based on
                                       setting of inqr
C
```

2.1.7. Function biniqr8 (Retrieving System-Dependent Parameters)

```
*deck,biniqr8
     function biniqr8 (nblk,key)
c *** primary function: get data about a block i/o buffer
c --- This routine is intended to be used in standalone programs.
c --- This routine should not be linked into the ANSYS program.

c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
c nblk (int,sc,in) - the block number for the inquiry
```

```
C
                                     or zero (see below)
                                   - key for information requested
С
      key
                (int,sc,in)
                       nblk = 0 - return information about system/file
C
С
                                key = 1 - return system block size
С
                                     = 2 - return number of integers per dp
                                     = 3 - return filename length
C
                                       5 = return integers per LONG
С
                       nblk > 0 - return information about this block
С
                                 key = 1 - return fortran unit number
C
С
                                     = 2 - return number of pages in file
                                     = 3 - return length of page (32 bit words)
C
                                     = 4 - return open status
С
С
                                           0 - file close
                                           1 - file open
C
                                     = 5 - return file format
С
                                           0 - internal format
С
                                           1 - external format
С
                                     = 6 - return read/write status
С
                                           0 - both read & write
С
                                           1 - read
C
                                           2 - write
C
                                     = 7 - return current position on file
C
С
                                     = 8 - return maximum length of file
                                             (in words)
С
                                     = 9 - return starting word for this page
C
                                             in buffer
С
                                     =10 - return base location
С
C
                                     =11 - return debug key
                                     =12 - return absolute (non-base) key
С
                                     =15 - return max record written
С
                                     =16 - return swap and record header key
C
                                     =17 - return precision key
С
   output arguments:
C
С
      binigr8
               (LONG, func, out)
                                  - the returned value of biniqr is based on
                                         setting of nblk and key
C
```

2.1.8. Function binset (Opening a Blocked Binary File or Initializing Paging Space)

```
*deck.binset
     function binset (nblk,nunit,ikeyrw,istart,paglen,npage,pname,
                       nchar, kext, Buffer4)
c *** primary function: initialize paging space for a blocked binary file.
С
                        binset should be used to open a blocked file
C
                        before binrd8 or binwrt8 are used. binclo should
                        be used to close the file.
c --- This routine is intended to be used in standalone programs.
c --- This routine should not be linked into the ANSYS program.
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
      nblk
             (int,sc,in)
                                  - block number (1 to BIO_MAXFILES max)
C
С
      nunit
                 (int,sc,in)
                                  - fortran unit number for the file
                                    (if 0, bit bucket)
С
      ikeyrw
                (int,sc,in)

    read/write flag

C
                                    = 0 - both read & write
С
                                    = 1 - read
С
С
                                       2 - write
С
                                    = 9 - read only
        NOTE: 0 may write, but the file length may not be extended and
C
                the file may or may not exist
С
              1 reads only, but the file protection must set set to "rw"
```

```
2 may extend the file length and the file is a new file
С
С
              9 reads only, but the file protection may be "r" only
                                  - starting location in buffer array
      istart
                (int,sc,in)
С
                                    usually 1 for nblk=1, paglen*npage+1
С
С
                                    for nblk=2,etc.
                                  - page length in integer*4 words for external
      paglen
С
                (int,sc,in)
                                    files
С
                                    paglen should always be a multiple of
С
                                    512 words for efficiency
С
С
                (int,sc,in)
                                  - number of pages (1 to BIO_MAXBLOCKS max)
                                  - name of the file
                (chr,ar(*),in)
С
      pname
С
      nchar
                (int,sc,in)
                                  - number of characters in the file name (not
С
                                    used)
                                  - no longer used, always external format
      kext.
                (int,sc,in)
C
               (i4, ar(*),inout) - work array for paging, should be
С
С
                                    dimensioned to paglen*npage*nblk (max)
  output arguments:
С
С
      binset (int,func,out)
                                  - error status
                                    = 0 - no error
C
                                    <>0 - error occurred
C
      Buffer4 (i4, ar(*),inout) - work array for paging
```

2.1.9. Subroutine bintfo (Defining Data for a Standard Mechanical APDL File Header)

```
*deck,bintfo
     subroutine bintfo (title,jobnam,units,code)
c *** primary function: set information necessary for binhed8
c --- This routine is intended to be used in standalone programs.
c --- This routine should not be linked into the ANSYS program.
C
c *** Notice - This file contains ANSYS Confidential information ***
C
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
C
C
  input arguments:
С
     variable (typ,siz,intent)
                                   description
               (chr*80,ar(2),in) - main title and 1st subtitle
С
     jobnam (chr*8,sc,in)
                               - jobname
C
С
     units
              (int,sc,in)
                                 - units
                                    = 0 - user defined units
С
                                    = 1 - SI
С
                                    = 2 - CSG
С
                                    = 3 - U.S. Customary, using feet
С
                                    = 4 - U.S. Customary, using inches
C
С
                                    = 5 - MKS
                                    = 6 - MPA
С
                                    = 7 - uMKS
С
С
      code
               (int,sc,in)
                                 - code defining 3rd party vendor
                                   (contact ANSYS, Inc. for code assignment)
C
С
c output arguments:
С
     none
С
```

2.1.10. Subroutine binhed8 (Writing the Standard Mechanical APDL File Header)

```
*deck.binhed8
     subroutine binhed8 (nblk,nunit,filposL,Buffer4)
c *** primary function:
                          put standard header on a binary file, all
                          permanent binary files should have this header
c *** secondary functions: return the first data position
c --- This routine is intended to be used in standalone programs.
c --- This routine should not be linked into the ANSYS program.
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
C
С
     nblk
              (int,sc,in)
                                 - block number of open binary file
                                   (as defined with subroutine binset)
С
                                 - the unit number for this file
     nunit.
              (int.sc.in)
C
С
     Buffer4 (int,ar(*),inout) - work array for paging, should be the
С
                                   same array as used in binset
  output arguments:
С
                                 - the position after the header
С
     filposL (LONG,sc,out)
     Buffer4 (int,ar(*),inout) - work array for paging
C
    ******* ANSYS standard header data description (100 words)
C
  loc
        no. words
                   contents
C
С
   1
           1
                    fortran unit number
    2
            2
                    file format
C
                    = 0 - old internal format
C
                    = -1 - external IEEE format
С
                    = 1 old external format
C
   3
           1
                    time in compact form (ie 130619 is 13:06:19)
С
С
    4
           1
                    date in compact form (ie 20051023 is 10/23/2005)
                    units
С
С
                    = 0 - user defined units
                    = 1 - SI
С
С
                     = 2 - CSG
                    = 3 - U.S. Customary, using feet
С
                    = 4 - U.S. Customary, using inches
C
                    = 5 - MKS
C
С
                    = 6 - MPA
С
                    = 7 - uMKS
                    User_Linked
С
c 10
                    revision in text format '10.0' (inexc4)
           1
c 11
                   date of revision release for this version
c 12
           3
                  machine identifier - 3 4-character strings
c 15
           2
                   (obsolete - see below) jobname - 2 4-character strings
                   product name - 2 4-character strings
С
  17
           2
c 19
           1
                   special version label - 1 4-character string
c 20
                   user name - 3 4-character strings
           3
c 23
                   machine identifier - 3 4-character strings
c 26
           1
                   system record size at file write
c 27
           1
                    maximum file length
C
  28
           1
                    maximum record number
c 31
                    jobname - 8 4-character strings
           8
c 41
           20
                   main title - 20 4-character strings
c 61
           20
                   first subtitle - 20 4-character strings
c 81
           1
                   file compression level
                    file sparsification key
С
  82
           1
С
  95
           1
                    split point of file
                    NOTE: Split files are not support by binlib!
c 96
           1
                   reserved for future use
c 97-98
                    LONGINT of file size at write
```

A version of binhed8 exists without the "8" suffix (binhed.F) that takes a regular integer for the third argument. This subroutine, therefore, cannot address large files where the file position exceeds 2**31. Use the binhed8.F version for any new program.

2.1.11. Subroutine binrd8 (Reading Data from a Buffered File)

```
*deck.binrd8
     subroutine binrd8 (nblk,LongLocL,leng,ivect,kbfint,Buffer4)
c ******* buffer read routine *******
c --- This routine is intended to be used in standalone programs.
c --- This routine should not be linked into the ANSYS program.
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
   nblk (int,sc,in)
                           - block number. see fd___(i.e. fdresu for results
C
                              file, etc.) include decks for more information.
C
   LongLocL(LONG,sc,inout) - location in integer*4 words of the startin
С
C
                              position on the file.
                           - number of words to read into ivect. (must be
         (int,sc,inout)
C
С
                              less or equal to dimension given to ivect in
C
                              the calling routine). if ivect is to be used
                              as integers, use as is. if ivect is to be
C
                              used for double precision numbers, it must be
С
                              increased by multiplying it by INTPDP.
C
                              if negative, skip record and do not return
С
                              data(results).
С
   Buffer4 (i4,ar(*),inout) - work array for paging, should be the
                              same array as used in binset
С
  output arguments:
C
   LongLocL(LONG,sc,inout) - location in integer*4 words of the current
                              position on the file. It is updated after
C
                              each read operation
С
  leng (int,sc,inout)
С
                           - tells you how many items it actually read(in
                              integer words).
C
С
                              if zero, end of file(error case)
  ivect (int,ar(*),out) - results (can be either integer or double
C
С
                              precision in the calling routine)
   kbfint (int,sc,out)
                            - key for type(used only for AUX2 dump)
С
                              = 0 double precision data
C
                              > 0 integer data(usually the same as leng)
C
  Buffer4 (i4,ar(*),inout) - work array for paging
```

Versions of binrd8/binwrt8 exist without the "8" suffix (binrd/binwrt) that take a regular integer for the second argument. These subroutines, therefore, cannot address large files where the file position exceeds 2**31. Use the binrd8/binwrt8 versions for any new programs.

2.1.12. Subroutine binwrt8 (Writing Data to a Buffered File)

```
c input arguments:
   nblk
          (int,sc,in)
                              - block number. see fd___(i.e. fdresu for results
C
                               file, etc.) include decks for more information.
C
С
   LongLocL(LONG, sc, inout)
                              - location in integer words of the starting
                               position on the file.
C
            (int,sc,in)
                               number of words to read from ivect. (must be
С
                                less or equal to dimension given to ivect in
С
                                the calling routine). if ivect is to be used
C
С
                                as integers, use as is. if ivect is to be
                                used for double precision numbers, it must be
С
C
                                increased by multiplying it by INTPDP.
С
   ivect
            (int,ar(*),in)
                              - data to be written onto the file(can be either
                                integer or double precision in the calling
C
С
                               routine)
С
   kbfint (int,sc,in)
                              - key for type(used only for AUX2 dump)
                               = 0 double precision data
С
                                > 0 integer data(usually the same as leng)
С
С
   Buffer4 (int,ar(*),inout) - work array for paging, should be the
                               same array as used in binset on this block
C
c output arguments:
   LongLocL(LONG,sc,inout)
                             - location in integer words of the current
С
                                position on the file. It is updated after
                               each write operation
C
            (int,ar(*),out) - vector containing record to be written
С
  Buffer4 (int,ar(*),inout) - work array for paging
```

Versions of binrd8/binwrt8 exist without the "8" suffix (binrd/binwrt) that take a regular integer for the second argument. These subroutines, therefore, cannot address large files where the file position exceeds 2**31. Use the binrd8/binwrt8 versions for any new programs.

2.1.13. Subroutine exinc4 (Decoding an Integer String into a Character String)

```
*deck,exinc4
      subroutine exinc4 (ichext,chin,n)
c primary function: decode externally formatted integer versions of 4-character
С
                    strings to plain 4-character strings (used to convert data
                    from externally formatted files to data for interally
С
                    formatted files)
C
c *** Notice - This file contains ANSYS Confidential information ***
C
  input arguments:
С
С
     ichext (int,ar(n),in)
                                - externally formatted integer form of
                                   4-character strings
C
               (int,sc,in)
                                 - number of strings to convert
С
C
С
  output arguments:
              (char,ar(n),out) - strings in character form
```

2.1.14. Subroutine inexc4 (Coding a Character String into an Integer String)

```
*deck,inexc4
subroutine inexc4 (chin,ichext,n)
c primary function: encode plain 4-character strings into externally formatted
c integer versions of 4-character strings (used to convert
c data from internally formatted files to data for
c externally formatted files)
```

```
c
c *** Notice - This file contains ANSYS Confidential information ***
c
c input arguments:
c chin (char,ar(n),in) - strings in character form
c n (int,sc,in) - number of strings to convert
c
c output arguments:
c ichext (int,ar(n),out) - externally formatted integer form of
c 4-character strings
```

2.1.15. Subroutine binclo (Closing or Deleting a Blocked Binary File)

```
*deck,binclo
     subroutine binclo (nblk,pstat,Buffer4)
c *** primary function: close a blocked file, every block/file opened with
                      binset should be closed with binclo
c *** secondary function: the file can be deleted by specifying 'D' in pstat
c --- This routine is intended to be used in standalone programs.
c --- This routine should not be linked into the ANSYS program.
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
                                - the block number to close
С
     nblk (int,sc,in)
С
                                   (as defined with subroutine binset)
    pstat
              (chr,sc,in)
                                 - keep or delete flag
C
C
                                  = 'K' - keep file
                                   = 'D' - delete file
С
    Buffer4 (int,ar(*),inout) - work array for paging, should be the
С
                                   same array as used in binset
c output arguments:
     Buffer4 (int,ar(*),inout) - work array for paging
```

2.1.16. Subroutine largeIntGet (Converting Two Integers into a Pointer)

```
*deck,largeIntGet
    function largeIntGet (small,large)

c primary function:    Convert two short integers into a long integer

c object/library: res

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
    c small (int,sc,in) - least significant part
    c large (int,sc,in) - most significant part

c output arguments:
    c largeIntGet (LONGINT,sc,out) - 64 bit integer
```

2.2. Demonstration Routines

The demonstration routines demonstrate several ways to use the binary file access routines provided with Mechanical APDL. The programs described below (all available on your distribution media; see Accessing Mechanical APDL Binary Files (p. 69) for their location) demonstrate other tasks that the binary access routines can do.

2.2.1. Program bintst (Demonstrates Dumping a Binary File and Copying It for Comparison Purposes)

The bintst program dumps a binary file with the name file.rst to the screen. It then takes that file, copies it to a new file, file2.rst, and dumps the new file to the screen for comparison purposes.

2.2.1.1. Common Variables:

Variable	Type, Size, Intent	Description
iout	int, sc, comm	The output unit number
intpdp	int, sc, comm	The number of integers per double precision word
lenfnm	int, sc, comm	The number of characters in the filename
reclng	int, sc, comm	The system record length

Note:

The bintst program is not part of the binlib.a library. It is included here only to aid you.

2.2.2. Subroutine bintrd (Demonstrates Printing a Dump of File Contents)

```
*deck,bintrd
     subroutine bintrd (pname)
  *** primary function: bin file dump utility
c *** Notice - This file contains ANSYS Confidential information ***
C
c Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
С
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
C
c input arguments:
     variable (typ,siz,intent) description
С
     pname (chr,sc,in)
                                  - name of binary file which is to
C
                                    be dumped to the screen
С
С
  output arguments:
С
C
     none.
С
  common variables:
С
              (int,sc,comm) - output unit number
(int,sc,comm) - number of integers per double precision word
С
     iout
C
     intpdp
                                - number of characters in the filename
С
     lenfnm
             (int,sc,comm)
С
    reclng (int,sc,comm)
                                 - system record length
C
                            NOTE: bintrd is not part of binlib.a. it is
C
```

```
c included only as an aid to users.
```

Note:

The bintrd routine and the bintwr routine described below are not part of binlib.a. This chapter includes it only to aid you. You can find the source for this routine on the product-distribution media.

Both subroutines require the following common:

```
COMMON/BINTCM/ IOUT,INTPDP,LENFNM,RECLNG
```

- Iout is the output unit number.
- Intpdp is the number of integers per double precision word.
- Lenfnm is the number of characters in the filename.
- Reclng is the system record length.

2.2.3. Subroutine bintwr (Demonstrates Copying Binary File Contents)

```
*deck,bintwr
     subroutine bintwr (pname, nname)
c *** primary function: bin file copy utility
c *** Notice - This file contains ANSYS Confidential information ***
С
c Copyright ANSYS. All Rights Reserved.
c *** ansys, inc.
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
С
C
c input arguments:
    variable (typ,siz,intent) description
C
    pname (chr,sc,in) - name of binary file which is to be copied
C
С
c output arguments:
    variable (typ,siz,intent) description
C
C
    nname (chr,sc,out) - name of new binary file which is a copy
                                 of pname
C
c common variables:
    iout (int,sc,comm) - output unit number intpdp (int,sc,comm) - number of integers per double precision word
С
C
    lenfnm (int,sc,comm) - number of characters in the filename
С
C
    reclng (int,sc,comm)
                                - system record length
C
                          NOTE: bintwr is not part of binlib.a. it is
С
С
                                included only as an aid to users.
C
```

2.2.4. Program ResRdDemo (Demonstrates Reading a Results File)

Program ResRdDemo demonstrates how to read a results file using the results file access routines (p. 90). The file must be named test.rst and the file contents are written to the screen.

This file resides in the subdirectory \Program Files\ANSYS Inc\V211\ANSYS\custom-ize\user (on Windows systems) or /ansys_inc/v211/ansys/customize/misc (on Linux systems).

2.2.5. Program ResWrDemo (Demonstrates Writing a Results File)

Program ResWrDemo demonstrates how to write a Mechanical APDL-readable results file. This file resides in the subdirectory \Program Files\ANSYS Inc\V211\ANSYS\customize\user (on Windows systems) or /ansys_inc/v211/ansys/customize/misc (on Linux systems).

2.3. Retrieving Data from the Results File

There are two methods for retrieving data from the results file:

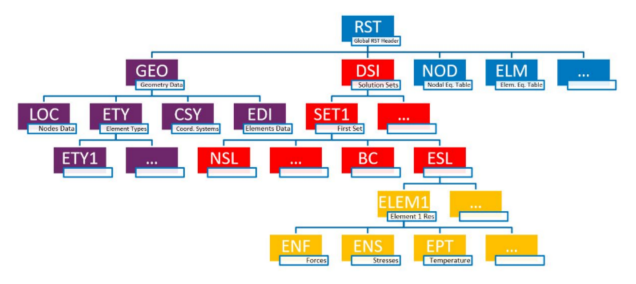
- 2.3.1. Results File Reader
- 2.3.2. Results File Access Routines

2.3.1. Results File Reader

The Mechanical APDL solver results are typically stored in a binary file using the format described in Description of the Results File (p. 5). This file can have an .rst, .rth, or .rmg extension, depending on the analysis type.

The results file format can be visualized as a tree of records.

Figure 2.1: Results File Structure



RST File Structure

Color code: blue - general records; purple - geometry; red - results sets; yellow - element results

The Results File Reader is a C++ object API that facilitates access to these records.

You can relate the above tree structure to the records described in Results File Format (p. 6). The below table shows typical examples.

Table 2.1: Finding Records in the Results File Description

Record Label GEO	Meaning	Sample Content from .rst File					
	Geometry data	c c 	GEO	i	1	80	Geometry data header(was 20 in 32 bit of the second of the
DSI	Data set index for solution sets	c c c c	DSI	i	1	2*resmax	Data sets index table. This record con the record pointers for the beginning each data set. The first resmax record the first 32 bits of the index, the se resmax records are the second 32 bits.
NOD	Nodal equivalence table	c c c	NOD	i	1	nnod	Nodal equivalence table. This table eq the number used for storage to the act node number (Back(i),i=1,nnod)
ELM	Element equivalence table	c c c	ELM	i	1	nelm	Element equivalence table. The program stores all element data in the numeric order that the SOLUTION processor solv elements. This table equates the order number used to the actual element numb

The following sections describe how to use the Results File Reader:

- 2.3.1.1. Compile and Link with the Result File Reader Code
- 2.3.1.2. Open an Existing Results File
- 2.3.1.3. Basic Concepts
- 2.3.1.4. Extract Nodes and Elements
- 2.3.1.5. Extract a Solution Vector
- 2.3.1.6. Example: Extract a Nodal Solution Vector
- 2.3.1.7. Example: Extract an Element Solution Vector
- 2.3.1.8. Example: Using the Results File Reader in a Standalone Program

2.3.1.1. Compile and Link with the Result File Reader Code

To compile your code based on the Results File Reader, you must add the path that contains the main C_RstFile.h include file and its dependencies, as shown below.

Windows:

C:\Program Files\ANSYS Inc\v211\ansys\customize\include

Linux:

/ansys_inc/v211/ansys/customize/include/

On Windows, you must add the -DWINNT preprocessor definition at compile time. On Linux, you must add -DLINUX.

To link your application, add the following libraries to your link process.

Windows:

```
C:\Program Files\ANSYS Inc\v211\ansys\custom\lib\winx64\libCadoeKernel.lib
C:\Program Files\ANSYS Inc\v211\ansys\custom\lib\winx64\libCadoeReaders.lib
```

Linux:

```
/ansys_inc/v211/ansys/lib/linx64/libCadoeKernel.so
/ansys_inc/v211/ansys/lib/linx64/libCadoeReaders.so
```

Note that these libraries have dependencies to the Intel MKL library.

2.3.1.2. Open an Existing Results File

The first step is to add this line at the beginning of your C++ code:

```
#include "C_RstFile.h"
```

Suppose that you have a results file named file.rst in your current directory. Use this line to open an existing file:

```
C_RstFile RstFile( "file.rst"); // File is opened. Ready to be parsed.
```

The file will automatically close when the object is deleted.

2.3.1.3. Basic Concepts

The following are basic concepts related to using the Results File Reader.

- 2.3.1.3.1. Vector Format Description
- 2.3.1.3.2. Get a Record in the Results File
- 2.3.1.3.3. Repeatedly Reading Records of the Results File

2.3.1.3.1. Vector Format Description

The Results File Reader offers the ability to get the values of a record of the results file via the CVEC<T> object, where T is the type of scalars of the record of interest (typically int, double, float or complex<double>).

You can create and allocate a CVEC object by typing:

```
CVEC<double> V(10); // Create and allocate a vector of 10 double values
```

The C_RstFile object allocates the vector using the correct size.

The size of a vector is accessible by calling the member function:

```
int size = V.Dim();
```

To access the values of the vector, get the pointer to the beginning of the vector:

```
double *values = V.Adr();
```

Or you can directly access the *i*th value:

```
int     i = 0;     // The first value
double val = V[i];
```

2.3.1.3.2. Get a Record in the Results File

Based on the tree representation shown in Figure 2.1: Results File Structure (p. 81), you can get any record from an opened results file just by specifying its location in the tree.

1. First, the global header of the results file can be accessed using a special call:

```
C_Record *RstHeader = RstFile.GetGlobalHeader(); // The main global RST File Header ("RST"
```

The results file header contains global information about what can be found in the current file (mesh, results, and so on).

To get the array of values from a C_Record object, use this call:

```
CVEC<int> VecHead; // VecHead is defined as an Vector of Integers.
RstFile.GetRecord( *RstHeader, VecHead); // Fills the VecHead vector
VecHead.Print(); // Print to screen the values of the vector
```

You can then access the values of this vector. Please refer to the C_RstFile::E_RstHeader C++ enum for a list of usable constants.

2. Next, from this global header you can get the C_Record object by giving the complete path to the record of interest. Here we access the nodal solution (NSL) of the first solution set:

```
C_Record *Block = RstHeader->find( "RST::DSI::SET1::NSL");
```

3. Then, depending on the type of values the record contains, you can get the values:

2.3.1.3.3. Repeatedly Reading Records of the Results File

Some sets of records are repeated in the file. To optimize the reading of such records, you can use the dedicated functions C_RstFile::GetFirstRec and C_RstFile::GetNextRec. They can be used for:

• The element descriptions:

RST::GEO::EID

The element type:

```
RST::GEO::ETY
```

• The element section data:

```
RST::GEO::SEC
```

• The coordinate system description:

```
RST::GEO::CSY
```

See Extract the Elements (p. 86) for examples of how to use these functions.

2.3.1.4. Extract Nodes and Elements

From the VecHead global header vector, you can get global information for the nodes.

· Maximum node number of the model:

```
C_RstFile::E_MAXN
```

· Actual number of nodes used in the solution phase:

```
C_RstFile::E_NNOD
```

Number of DOFs per node:

```
C_RstFile::E_NUMDOF
```

2.3.1.4.1. Extract the Nodes

Node locations are stored in the record "LOC", under the geometry section of the tree:

```
c LOC dp nnod 7 (64 bit version)
c Node,X,Y,Z,THXY,THYZ,THZX for each node
c Nodes are in node number order
```

To get the entire vector of node numbers and coordinates, you can use this direct call:

2.3.1.4.2. Extract the Elements

From the Global header vector, you can get global information for the elements.

· Maximum element number of the model:

```
C_RstFile::E_MAXE
```

· Actual number of elements used in the solution phase:

```
C_RstFile::E_NELM
```

Element Description Record

Element descriptions are stored in the record "EID", in the geometry section of the tree:

```
Element descriptions index table. This
    ETD
                           nelm
C
С
                                    record contains the record pointers for each
С
                                    element description. (LONGINT (2*nelm) for
C
                                    64 bit version, relative to ptrEIDL).
                                    The order of the elements is the same as
C
                                    the order in the element equivalence table.
C
                   nelm 10+nodelm Element descriptions. Each of these records
С
C
                                    is pointed to by a record pointer given in
С
                                    the record labeled EID. The length of these
С
                                    records varies for each element (actual
                                    length is returned from routine BINRD8).
C
С
                                    nodelm shown here is the number of nodes for
                                    this element. Its value is defined in the
C
                                    element type description record.
C
```

You can, for instance, get repeatedly all the element descriptions by calling the functions C_RstFile::GetFirstRec and C_RstFile::GetNextRec:

```
CVEC<int> ElemDesc;
int iel = RstFile.GetFirstRec( "RST::GEO::EID", ElemDesc, 0, true);
while (iel != -1)
    {
      iel = RstFile.GetNextRec( ElemDes);
    }
```

Element Type Record

Element type descriptions are stored in the record "ETY":

```
c ETY i 1 maxety The element types index table. This record contains record pointers for each element type description. (Relative to ptrETYPL for 64 bit version)
```

In the same way as shown above for element descriptions, you can loop over the element types, calling the same set of functions.

```
CVEC<int> ElemType;
int iel = RstFile.GetFirstRec( "RST::GEO::ETY", ElemType, 0, true);
while (iel != -1)
    {
    iel = RstFile.GetNextRec( ElemType);
}
```

2.3.1.5. Extract a Solution Vector

To get a given nodal or element solution in a given solution set, you can call the C_RstFile::Get-Solution function.

```
CVEC<T> *V = RstFile.GetSolution(int iset, string IdSol, CVEC<T> *Sol, int NumEntity);
```

where:

iset is the solution set number.

IdSol is the solution record ID (for example, "NSL", "RF", "ENS", and so on).

Sol is the vector where the values are to be stored. If it is not allocated, the function allocates it and returns.

NumEntity is the node or element number. Default = All_REC, meaning all entities in the same vector are requested.

You can access the vector properties using this:

```
int Size = SolNod.Dim(); // To get the size of the output vector
double val = SolNod[ii]; // To access the values of the vector
```

2.3.1.6. Example: Extract a Nodal Solution Vector

This example demonstrates reading the nodal solution corresponding to the first solution set.

2.3.1.7. Example: Extract an Element Solution Vector

This example demonstrates reading the element nodal stresses corresponding to the first solution set and the first element.

Given an elementary result, we can also loop over all elements where this result is stored using the C_RstFile::GetFirstLocalSolution and C_RstFile::GetNextLocalSolution functions.

```
int
              iSet = 1; // ===== Solution set number
                                 // ===== Solution Id we want to get
                IdSol = "ENS";
string
CVEC<double>
                                   // ===== The vector to fill with the solution
               EnsVector;
               NumEntity = 1;
                                  // ===== The first element number
NumEntity = RstFile.
GetFirstLocalSolution( iSet, IdSol, EnsVector, NumEntity);
While ( NumEntity > 0)
      // EnsVector contains nodal stresses of the element #NumEntity
     NumEntity
     = RstFile.
     GetNextLocalSolution( EnsVector, NextEntity);
```

2.3.1.8. Example: Using the Results File Reader in a Standalone Program

```
* @file TestRstReader.cpp
* @brief This source file shows the way we can open an RST File,
* and extract data using the C++ C_RstFile Object.
* /
#include <stdio.h>
#include "C_RstFile.h"
main(int argc, char *argv[])
char *FileName = argv[1];
// ===== Open the RST File :
cout << "\n Open the RST File : " << FileName << endl;</pre>
C_Record *RstHeader = RstFile.GetGlobalHeader(); //
                                                  Handle on the Global header
cout << "\n Ansys Rev. : " << RstFile.AnsRev() << endl;</pre>
// ===== Get global numbers for this model
int maxn = RstFile.RstHeader(C_RstFile::E_MAXN); // Maximum node number
int nnod = RstFile.RstHeader(C_RstFile::E_NNOD); // Number of nodes
int maxe = RstFile.RstHeader(C_RstFile::E_MAXE); // Maximum element number
int nelm = RstFile.RstHeader(C_RstFile::E_NELM); // Number of elements
cout << "\n Number of Nodes : " << nnod;</pre>
cout << "\n Max Node Number : " << maxn << endl;</pre>
cout << "\n Number of Elements : " << nelm;</pre>
cout << "\n Max Elt Number : " << maxe << endl;</pre>
// ===== Extract Nodes Data
// Node,X,Y,Z,THXY,THYZ,THZX for each node. Nodes are in node number order
C_Record *Block = NULL;
Block = RstHeader->find("RST::GEO::LOC"); // Get the pointer to the LOC Record
```

```
CVEC<> GeoNod;
RstFile.GetRecord(*Block, GeoNod, ALL_REC); // Fills the GeoNod Array
cout << "\n Global Nodes Vector, Size = : " << GeoNod.Dim();</pre>
// The same data, but for a single node:
int NumNod(1);
CVEC<> GeoNod i;
RstFile.GetRecord(*Block, GeoNod_i, NumNod); // We only read the data for the 1st node
cout << "\n Single Nodes Vector, Size = : " << GeoNod_i.Dim() << endl;</pre>
// ===== Extract Elements Data
// ===== Mapping Elt Index <-> Elt Number
Block = RstHeader->find("RST::ELM"); // Get the pointer to the ELM Record
CVEC<int> Elm;
RstFile.GetRecord(*Block, Elm); // Fills the Elm Array
cout << "\n Element Equivalence Table Size : " << Elm.Dim() << endl;</pre>
// ===== Get the Geometry Header
Block = RstHeader->find("RST::GEO"); // Get the pointer to the GEO Record
CVEC<int> GeoHeader;
RstFile.GetRecord(*Block, GeoHeader); // Fills the GeoHeader Array
// This GeoHeader contains information such as the maximum element type reference number in the model:
int maxety = GeoHeader[C_RstFile::E_GMAXETY];
// For each element type, we can get the Type description
cout << "\n Extract All Element Types Descriptions .... " << endl;</pre>
CVEC<int> ElemEty;
int ityp = RstFile.GetFirstRec("RST::GEO::ETY", ElemEty, 0, true);
while (ityp != -1)
cout << " Got Description for Element Type : " << ElemEty[1] << endl;</pre>
ityp = RstFile.GetNextRec(ElemEty);
// For each element, we extract the description associated to
cout << "\n Extract All Element Descriptions .... \n";</pre>
CVEC<int> ElemDes;
int iel = RstFile.GetFirstRec("RST::GEO::EID", ElemDes, 0, true);
while (iel !=-1)
int imat = ElemDes[0],
 itype = ElemDes[1];
cout << "Element " << iel << ": Material = " << imat << " , Element Type Number = " << itype << endl;
iel = RstFile.GetNextRec(ElemDes);
cout << "\n
                       Element " << nelm << " OK" << endl;</pre>
// ===== We get the number of solution sets
int NbSet = RstFile.RstHeader(C_RstFile::E_NSETS);
```

```
cout << "\n Number of Result Sets : " << NbSet << endl;</pre>
// Get the Nodal Solution of the 1st solution set
     iset(1); // 1st solution set
string IdSol = "NSL"; // "NSL" means Nodal Solution
CVEC<> SolNod; // This vector will contain the Nodal Solution
RstFile.GetSolution(iset, IdSol, &SolNod);
cout << "Nodal Solution vector : size = " << SolNod.Dim() << endl;</pre>
// Get the Nodal Stresses for the 1st Element, for the 1st solution set
IdSol = "ENS";
int NumElem = 1;
int Size(0);
CVEC<> EnsElem;
try {
RstFile.GetSolution(iset, IdSol, &EnsElem, NumElem);
catch (C_CadoeException e) {
if (e.getExceptionCode() == E_ResultNotAvailable)
  cerr << "This result " << IdSol << " is not available\n";</pre>
Size = EnsElem.Dim();
cout << "\n Size of the ENS vector for the 1st Element = " << Size << endl;</pre>
// Loop to get all the Element Nodal Stresses, for the 1st Solution Set
NumElem = RstFile.GetFirstLocalSolution( iset, IdSol, EnsElem, 1); // 1 means we start at the 1st Element
if (NumElem == -3) cerr << "No Elementary results stored in this Result File\n";
if (NumElem == -2) cerr << "this result is not stored for this Element " << NumElem << "in this Solution Set\n
while(NumElem > 0)
cout << "For Element #" << NumElem << " Norm of the Nodal Stress Vector is : " << EnsElem.Nrm() << endl;
NumElem = RstFile.GetNextLocalSolution( EnsElem);
return;
```

2.3.2. Results File Access Routines

The method of accessing the results file described in this section is presented as an alternative to the Results File Reader (p. 81).

You can use the low-level routines described in Accessing Mechanical APDL Binary Files (p. 69) to retrieve data from the results file. Or you can use the routines described below that retrieve data specific to the format of the results file.

These files reside in the subdirectory \Program Files\ANSYS Inc\V211\ANSYS\custom-ize\user (on Windows systems) or /ansys_inc/v211/ansys/customize/misc (on Linux)

systems). See Access Routines for Results Files (p. 69) for information on compiling and linking these routines.

2.3.2.1. Overview of the Routines

For each data record in the results file, routines exist that:

- Read the record index and allocate space for the data. These are named ResRdrecordBegin, where record is a descriptive name of the record, for example, ResRdNodeBegin
- Read the data itself. These are named ResRdrecord, for example, ResRdNode
- Deallocate space for the data. These are named ResRdrecordEnd, for example, ResRd-NodeEnd

Below is a complete listing of all the routines with the indentation indicating the required nested calling sequence:

```
function ResRdBegin (Nunit,Lunit,Fname,ncFname,Title,JobName,Units,NumDOF,DOF,UserCode,MaxNode,NumNode,MaxElem,
 subroutine ResRdGeomBegin (MaxType,MaxReal,MaxCsys,nXYZ)
    subroutine ResRdTypeBegin (NumType, TypeLen)
     function ResRdType (itype,ielc,TypeNo)
    subroutine ResRdTypeEnd
    subroutine ResRdRealBegin (NumReal, NumPerReal, RealLen)
      function ResRdReal (iReal, Rcon, RealNo)
    subroutine ResRdRealEnd
    subroutine ResRdCsysBegin (NumCsys,CsysLen)
     function ResRdCsys (iCsys,CsysNo)
    subroutine ResRdCsysEnd
    subroutine ResRdNodeBegin
     function ResRdNode (iNode, xyzang)
    subroutine ResRdNodeEnd
    subroutine ResRdElemBegin
      function ResRdElem (iElem, nodes, ElemData)
    subroutine ResRdElemEnd
 subroutine ResRdGeomEnd
 subroutine ResRdSectMatBegin (MaxSect, MaxMat)
    subroutine ResRdSectBegin (NumSect,NumPerSect,SectLen)
      function ResRdSect (iSect,SecData,SectNo)
    subroutine ResRdSectEnd
    subroutine ResRdMatBegin (NumMat,NumPerMat)
      function ResRdMat (iMat,iprop,MatData)
    subroutine ResRdMatEnd
 subroutine ResRdSectMatEnd
 function ResRdSolBegin (key, lstep, substep, ncumit, kcmplx, time, Title, DofLab)
    subroutine ResRdDispBegin
      function ResRdDisp (node,Disp)
    subroutine ResRdDispEnd
    subroutine ResRdRforBegin (nRForce)
      function ResRdRfor (node,idof,value)
    subroutine ResRdRforEnd
    subroutine ResRdBCBegin (BCHeader)
      subroutine ResRdFixBegin (BCHeader,nFixed)
        function ResRdFix (node,idof,value)
      subroutine ResRdFixEnd
      subroutine ResRdForcBegin (BCHeader,nForces)
        function ResRdForc (node, idof, value)
      subroutine ResRdForcEnd
    subroutine ResRdBCEnd
    subroutine ResRdEresBegin
      function ResRdEstrBegin (iElem)
        function ResRdEstr (iStr,Str)
      subroutine ResRdEstrEnd
    subroutine ResRdEresEnd
```

```
subroutine ResRdSolEnd
subroutine ResRdEnd
```

These routines are contained in the file ResRd. F. See the demonstration routine ResRd-Demo.F (p. 80) on the distribution medium for an example of the usage of these routines.

The memory allocation scheme is described in Memory Management Routines (p. 272) in Part 2: Guide to User-Programmable Features (p. 125).

The following sections describe the data-reading routines. See the file ResRd.F and its corresponding include deck ResRd.inc for listings of the corresponding Begin/End routines.

2.3.2.2. ResRdBegin (Opening the File and Retrieving Global Information)

```
*deck,ResRdBegin
      function ResRdBegin (Nunit, Lunit, Fname, ncFname, Title, JobName,
                              Units, NumDOF, DOF, UserCode,
                              MaxNode, NumNode, MaxElem, NumElem,
     Х
                              MaxResultSet, NumResultSet)
     x
c primary function:
                        Open result file and return global information
c object/library: ResRd
c input arguments:
     Nunit (int,sc,in) - Fortran Unit number for file (ANSYS uses 12)
Lunit (int,sc,in) - Current print output unit (usually 6 <STDOUT>)
C
С
      Fname
                (ch*(ncFname),sc,in) - The name (with extension) for the file
С
      ncFname (int,sc,in) - Number of characters in Fname
С
c output arguments:
    Title (ch*80,ar(2),out) - Title and First subtitle
С
      JobName (ch*32,sc,out) - Jobname from file Units (int,sc,out) - unit system
С
      Units (int,sc,out)
C
С
                                        = 0 - user defined units
С
                                         = 1 - SI
                                         = 2 - CSG
C
С
                                         = 3 - U.S. Customary, using feet
                                         = 4 - U.S. Customary, using inches
С
                                         = 5 - MKS
C
                                         = 6 - MPA
С
                                        = 7 - uMKS
С
               (int,sc,out)
                                   - Number of DOF per node
      NumDOF
С
     DOF (int,ar(*),out) - The DOFs per node
UserCode (int,sc,out) - Code for this application
MaxNode (int,sc,out) - Maximum node number used
С
C
С
C
    NumNode (int,sc,out)
                                   - Number of nodes attached to elements
    MaxElem (int,sc,out) - Maximum element number used
NumElem (int,sc,out) - Number of elements used
C
      MaxResultSet (int,sc,out) - Maximum number of result sets (usually 1000)
С
C
      NumResultSet (int,sc,out) - Number of result sets on file
     ResRdBegin (int,sc,out) - 0, successful other, error in file open
```

2.3.2.3. ResRdGeomBegin (Retrieving Global Geometry Information)

```
*deck,ResRdGeomBegin subroutine ResRdGeomBegin (MaxType, MaxReal, MaxCsys, nXYZ)
c primary function: Read Geometry Header Record
c object/library: ResRd
c input arguments: none
```

```
c output arguments:
c MaxType (int,sc,out) - Maximum element type
c MaxReal (int,sc,out) - Maximum real constant set number
c MaxCsys (int,sc,out) - Maximum coordinate system number
c nXYZ (int,sc,out) - number of nodes with coordinates
```

2.3.2.4. ResRdType (Retrieving Element Types)

```
*deck,ResRdType function ResRdType (itype,ielc,TypeNo)
c primary function: Read an element type record

c object/library: ResRd

c input arguments:
c itype (int,sc,in) - Element type number

c output arguments: none
c ielc (int,ar(IELCSZ),out) - Element characteristics
c TypeNo (int,sc,out) - External Element Type number
c ResRdType (int,sc,out - number of words read
```

2.3.2.5. ResRdReal (Retrieving Real Constants)

```
*deck,ResRdReal function ResRdReal (iReal,Rcon,RealNo)
c primary function: Read real constant record

c object/library: ResRd

c input arguments:
c iReal (int,sc,in) - Real set number

c output arguments: none
c Rcon (dp,ar(ResRdReal),out) - Real Constants
c RealNo (int,sc,out) - External Real Constant number
c ResRdReal (int,sc,out) - Number of real constants in set
```

2.3.2.6. ResRdCsys (Retrieving Coordinate Systems)

```
*deck,ResRdCsys
function ResRdCsys (iCsys,Csys,CsysNo)
c primary function: Read a coordinate system record

c object/library: ResRd

c input arguments:
c iCsys (int,sc,in) - Coordinate system number

c output arguments:
c Csys (dp,ar(ResRdCsys),out) - Coordinate system description
c CsysNo (int,sc,out) - External Coordinate system number
c ResRdCsys (int,sc,out) - Number of values
```

2.3.2.7. ResRdNode (Retrieving Nodal Coordinates)

2.3.2.8. ResRdElem (Retrieving Elements)

```
*deck,ResRdElem
     function ResRdElem (iElem, nodes, ElemData)
c primary function: Read an element
c object/library: ResRd
c input arguments:
     iElem (int,sc,in) - The element number
c output arguments:
   ResRdElem(int,sc,out)
                                - Number of nodes
С
     nodes (int,ar(n),out) - Element nodes
С
     ElemData (int,ar(10),out) - Element information
C
С
                                    mat - material reference number
С
                                    type - element type number
                                    real - real constant reference number
C
С
                                    secnum - section number
                                    esys - element coordinate system
С
                                    death - death flag
C
                                             = 0 - alive
                                             = 1 - dead
С
                                    solidm - solid model reference
C
                                    shape - coded shape key
elnum - element number
С
C
                                    pexcl - P-Method exclude key
```

2.3.2.9. ResRdSectMatBegin (Retrieving Global Section and Material Information)

```
*deck,ResRdSectMatBegin subroutine ResRdSectMatBegin (MaxSect, MaxMat)
c primary function: Read maximum section and material number from the Geometry Header Record
c object/library: ResRd
c input arguments: none
c output arguments:
c MaxSect (int,sc,out) - Maximum section number
c MaxMat (int,sc,out) - Maximum material number
```

2.3.2.10. ResRdSect (Retrieving Section Data)

```
*deck,ResRdSect function ResRdSect (iSect,SecData,SectNo)
c primary function: Read section record

c object/library: ResRd

c input arguments:
c iSect (int,sc,in) - Section set number

c output arguments:
c SecData (dp,ar(ResRdSect),out) - Section data
c SectNo (int,sc,out) - Exteral Section number

c ResRdSect (int,sc,out) - Number of section data in set
```

2.3.2.11. ResRdMat (Retrieving Material Data)

```
*deck,ResRdMat
     function ResRdMat (iMat,iprop,MatData)
c primary function: Read material record
c object/library: ResRd
c input arguments:
    iMat (int,sc,in)
                                  - Material set number
C
     iprop (int,sc,in)
                                  - Property reference number
C
                                     See mpingr for details
C
c output arguments:
     MatData (dp,ar(ResRdMat),out) - Material data for type iprop
C
     ResRdMat (int,sc,out) - Number of material data in set
```

See Function mpinqr (Getting Information About a Material Property) (p. 294) for details on the property reference number (iprop).

2.3.2.12. ResRdSolBegin (Retrieving Result Set Location)

```
*deck,ResRdSolBegin
     function ResRdSolBegin (key, lstep, substep, ncumit, kcmplx, time,
                              Title, DofLab)
c primary function: Read the solution header records
c object/library: ResRd
c input arguments:
                                 - 0, find by set number
    key (int,sc,in)
C
                                  1, find by lstep/substep
C
С
                                   2, find by ncumit
С
                                  3, find by time
     lstep (int,sc,in/out) - Load step number
С
                                   if key=0, this is the set number
С
     substep (int,sc,in/out) - Substep of this load step
C
     ncumit (int,sc,in/out) - Cumulative iteration number
C
     kcmplx (int,sc,in) - 0, Real solution 1,
time (dp,sc,in/out) - Current solution time
                                - 0, Real solution 1, Imaginary solution
C
С
c output arguments:
С
     Title (ch*80,ar(5),out) - Title and 4 subtitles
     DofLab (ch*4,ar(nDOF),out) - Labels for DOFs
```

```
c ResRdSolBegin (int,sc,out) - 0, requested solution set found c 1, not found
```

2.3.2.13. ResRdDisp (Retrieving Nodal Solution)

```
*deck,ResRdDisp
    function ResRdDisp (node,Disp)
c primary function: Retrieve a nodal displacement

c object/library: ResRd

c input arguments:
c node (int,sc,in) - Node number

c output arguments: none
c Disp (dp,ar(nDOF),out) - Displacements
c ResRdDisp(int,sc,out) - Number of displacements
```

2.3.2.14. ResRdRfor (Retrieving Reaction Solution)

```
*deck,ResRdRfor
function ResRdRfor (node,idof,value)
c primary function: Retrieve a reaction force

c object/library: ResRd

c input arguments:
c node (int,sc,in) - External node number
c idof (int,sc,in) - Internal dof number

c output arguments:
c value (dp,sc,in) - Value of reaction force
c ResRdRfor (int,sc,out) - Number of returned values (0 or 1)
```

2.3.2.15. ResRdFix (Retrieving Applied Nodal Constraints)

```
*deck,ResRdFix function ResRdFix (node,idof,value)
c primary function: Retrieve a constraint value

c object/library: ResRd

c input arguments:
c node (int,sc,in) - External node number
c idof (int,sc,in) - Internal dof number

c output arguments:
c value (dp,ar(4),in) - Real,Imag, RealOld,ImagOld
c ResRdFix (int,sc,out) - Number of returned values (0 or 4)
```

2.3.2.16. ResRdForc (Retrieving Applied Nodal Loads Solution)

```
*deck,ResRdForc function ResRdForc (node,idof,value)
c primary function: Retrieve an applied force value
```

```
c object/library: ResRd

c input arguments:
c node (int,sc,in) - External node number
c idof (int,sc,in) - Internal dof number

c output arguments:
c value (dp,ar(4),in) - Real,Imag, RealOld,ImagOld
c ResRdForc (int,sc,out) - Number of returned values (0 or 4)
```

2.3.2.17. ResRdEstr (Retrieving Element Solutions)

```
*deck,ResRdEstr function ResRdEstr (iStr,Str)
c primary function: Get an element's results

c object/library: ResRd

c input arguments:
c iStr (int,sc,in) - element record number (1-NUMELEDATASETS)

c output arguments:
c ResRdEstr (int,sc,out) - Number of element values
c Str (dp,ar(nStr),out) - element values
```

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Chapter 3: The CDWRITE (CDB) File Format

This chapter discusses how to write a coded database file, <code>Jobname.CDB</code>, that can be used to export a model from Mechanical APDL into another application. The <code>Jobname.cdb</code> file contains model data in terms of Mechanical APDL input commands.

The following topics are discussed:

- 3.1. Using the CDWRITE Command
- 3.2. Coded Database File Commands

3.1. Using the CDWRITE Command

To export a model from Mechanical APDL to another application, use menu path **Main Menu> Preprocessor> Archive Model> Write** or the **CDWRITE** command within the general preprocessor, PREP7. This produces a coded database file called Jobname.cdb. You specify the jobname using **Utility Menu> File> Change Jobname** or the **/FILNAME** command. If you supply no jobname, Mechanical APDL uses the default name "file".

The Jobname.cdb file contains selected geometry (nodes and elements), load items, and other model data in terms of Mechanical APDL input commands. (For a complete list of data in the file, see the **CD-WRITE** description in the *Command Reference*.) You can convert this information to a format compatible with the program into which you are importing it. The next few pages describe special considerations and commands you may need to do this conversion.

Note:

Files created by the **CDWRITE** command have the active coordinate system set to Cartesian (**CSYS**, 0).

Mechanical APDL may create parameters in the **CDWRITE** file that start with an underscore (_), usually an "_z." Such parameters are for Mechanical APDL internal use and pass information to the Mechanical APDL GUI.

3.1.1. Customizing Degree of Freedom Labels: the /DFLAB Command

Mechanical APDL uses a set of default labels for the degrees of freedom. You use these labels when entering boundary conditions, or Mechanical APDL uses the labels when writing the Jobname.cdb file.

You can change the labels to reflect the degrees of freedom of the other program by issuing the command /DFLAB. If you are customizing the degree-of-freedom labels, /DFLAB must be the first command you enter within Mechanical APDL. You may want to include the command in your START. ANS file. You can use /DFLAB only at the Begin processing level.

/DFLAB assigns or reassigns the "displacement" and "force" labels in the Mechanical APDL degree-of-freedom list. For example, DOF number 1 is predefined to have a displacement label of UX and a force label of FX, but you can assign new labels to this degree of freedom by issuing **/DFLAB**. Changing predefined labels generates a warning message.

The format for the /DFLAB command is:

/DFLAB, NDOF, LabD, LabF

NDOF

Mechanical APDL degree-of-freedom number (1 to 32)

LabD

Displacement degree of freedom label to be assigned (up to four characters)

LabF

Force label to be assigned (up to four characters)

You can also use /**DFLAB** to assign labels to spare degree-of-freedom numbers. Spare displacement and force labels are from 27 to 32. All other degree-of-freedom numbers are predefined, as follows:

DOF Number	Corresponding Displacement Label	Corresponding Force Label
1	UX	FX
2	UY	FY
3	UZ	FZ
4	ROTX	MX
5	ROTY	MY
6	ROTZ	MZ
7	AX	CSGX
8	AY	CSGY
9	AZ	CSGZ
10	VX	VFX
11	VY	VFY
12	VZ	VFZ
13	GFV1	GFL1
14	GFV2	GFL2
15	GFV3	PFFL
16	WARP	ВМОМ
17	CONC	RATE
18	HDSP	DVOL
19	PRES	FLOW
20	TEMP	HEAT

DOF Number	Corresponding Displacement Label	Corresponding Force Label
21	VOLT	AMPS
22	MAG	FLUX
23	ENKE	NPKE
24	ENDS	NPDS
25	EMF	CURT
26	CURR	VLTG

3.2. Coded Database File Commands

In the coded database file Jobname. CDB, most Mechanical APDL commands have the same format they have elsewhere. (See the *Command Reference* for command-specific information.) However, the format for some commands differs slightly in the Jobname. CDB file. The format for these commands is described below. Commands that use an unblocked format include the label UNBL in one of the command fields.

The **CDWRITE** command has an UNBLOCKED and a BLOCKED option. The UNBLOCKED option writes all data out in command format. The default BLOCKED option writes certain data items in a fixed format, including those that could potentially contain large amounts of data, such as nodal data.

3.2.1. BFBLOCK Command

BFBLOCK defines a block of nodal body-force loads. This is the recommended method for inputting nodal body-force loads into the Mechanical APDL database. The command syntax is:

BFBLOCK, NUMFIELD, Lab, NDMAX, NDSEL, TAB, -, MESHFLAG Format

NUMFIELD

The number of fields in the blocked format.

Lab

The body-force load label used to describe the block data. (For a list of load labels, see BF.)

NDMAX

The maximum node number defined.

NDSEL

The number of selected nodes.

Reserved.

MESHFLAG

When using nonlinear adaptivity in a linear analysis (**NLGEOM**,OFF), specifies how to apply nodal body-force loading on the mesh. Valid only when Lab = HGEN or TEMP.

- 0 Nodal body-force loading occurs on the current mesh (default).
- 1 Nodal body-force loading occurs on the initial mesh for nonlinear adaptivity.

Format

Data descriptors defining the format.

The format of the body-force load block is as follows:

- Field 1 Node number.
- Fields 2-7 Body-force load values to apply to the node. The number of fields is dependent on the body-force load label. (See **BF**.)

The final line of the block format is always a BF command with -1 for the node number.

The following example shows a typical **BFBLOCK** formatted set of body-force loads (using non-tabular input) that define a temperature load (TEMP).

```
BFBLOCK, 2, TEMP,
                         97,
                                     97,0
(i9,6(pg16.9))
                300.000000
        1
        2
                300.000000
                300.000000
        3
       95
                300.000000
       96
                300.000000
       97
                300.000000
BF, end, LOC,
```

For an example using tabular input, see the **BFEBLOCK** command (p. 102).

In the GUI, the **BFBLOCK** command must be contained in an externally prepared file and read into Mechanical APDL (via **CDREAD**, /INPUT, or other commands).

The **BFBLOCK** command is not valid in a ***DO** loop.

3.2.2. BFEBLOCK Command

BFEBLOCK defines a block of element body-force loads. This is the recommended method for inputting element body-force loads into the Mechanical APDL database. The command syntax is:

```
BFEBLOCK, NUMFIELD, Lab, ELMAX, ELSEL, TAB
Format
```

NUMFIELD

The number of fields in the blocked format.

Lab

The body-force load label used to describe the block data. (See BFE for a list of load labels.)

ELMAX

The maximum element number defined.

ELSEL

The number of selected elements.

TAB

Key for tabular input:

- 0 Non-tabular input.
- 1 Tabular input.

Format

Data descriptors defining the format.

The format of the body-force load block is as follows:

- · Field 1 Element number.
- Field 2 Starting location for entering data.
- Fields 3 Body load value for the specified starting location. (See **BFE** for information about starting location and associated body-force load values.)

The final line of the block format is always a BFE command with -1 for the element number.

The following example shows a typical **BFEBLOCK** formatted set of body-force loads (using tabular input) that define a temperature load (TEMP).

```
BFEBLOCK, 3, TEMP,
                         108,
                                     108,1
(2i9,a)
                   1
                              %BODYFORCE%
         2
                              %BODYFORCE%
                   1
         3
                   1
                              %BODYFORCE%
       106
                   1
                              %BODYFORCE%
       107
                   1
                              %BODYFORCE%
       108
                   1
                              %BODYFORCE%
                    -1,
BFE, end, LOC,
```

The following example shows a typical **BFEBLOCK** formatted set of body-force loads (using non-tabular input) that define a force load (FORC) with complex input.

In the GUI, the **BFEBLOCK** command must be contained in an externally prepared file and read into Mechanical APDL (via **CDREAD**, /**INPUT**, or other commands).

The **BFEBLOCK** command is not valid in a ***DO** loop.

3.2.3. CE Command

CE defines the constant term in a constraint equation. The command format in Jobname. CDB is:

```
{\tt CE,UNBL,Type,LENGTH,NCE,CONST}
```

Type

The type of data to be defined. DEFI is the valid label.

LENGTH

The total number of variable terms in the constraint equation.

NCE

The constraint equation reference number.

CONST

The constant term of the equation.

Another version of the **CE** command defines the variable terms in a constraint equation. You must issue this version of the command after the **CE** command described above. This command repeats until all terms are defined.

The alternate format for the **CE** command is:

```
CE, UNBL, Type, N1, Dlab1, C1, N2, Dlab2, C2
```

Type

The type of data to be defined. NODE is the valid label.

N1

The node number of the next term.

Dlab1

The degree-of-freedom label of N1.

C1

The coefficient of N1.

N2

The node number of the next term.

Dlab2

The degree-of-freedom label of N2.

C2

The coefficient of N2.

3.2.4. CP Command

CP defines a coupled-node set. You repeat the command until all nodes are defined. The command format in Johname. CDB is:

```
CP, UNBL, LENGTH, NCP, Dlab, N1, N2, N3, N4, N5, N6, N7
```

LENGTH

The total number of nodes in the coupled set

NCP

The coupled node reference number

Dlab

The degree of freedom label for the set

N1,N2,N3,N4,N5,N6,N7

The next seven node numbers in the coupled set

3.2.5. CMBLOCK Command

CMBLOCK defines the entities contained in a node or element component. The command format in Jobname. CDB is:

```
CMBLOCK,Cname,Entity,NUMITEMS,,,,,KOPT
Format
```

Cname

Eight character component name.

Entity

Label identifying the type of component (NODE or ELEMENT).

NUMITEMS

Number of items written.

--

Reserved for future use.

_ -

Reserved for future use.

_ _

Reserved for future use.

__

Reserved for future use.

KOPT

Controls how element component contents are updated during nonlinear mesh adaptivity analysis:

- 0 Component is not updated during remeshing and therefore contains only initial mesh elements (default).
- 1 Component is updated during remeshing to contain the updated elements.

This argument is valid only for nonlinear mesh adaptivity analysis with Entity = ELEM, and for solid element components only. This argument does not support NLAD-ETCHG analysis.

Format

Data descriptors defining the format. For **CMBLOCK** this is always (8i10).

The items contained in this component are written at 10 items per line. Additional lines are repeated as needed until all NumItems are defined. If one of the items is less than zero, then the entities from the item previous to this one (inclusive) are part of the component.

The **CMBLOCK** command is not valid in a ***DO** loop.

3.2.6. CYCLIC Command

CYCLIC,CDWR defines the input and output of a cyclic symmetry analysis. The syntax is:

```
CYCLIC, CDWR, Value1, Value2, Value3, ...
```

The following describes the values written to the . CDB file for cyclic options CYCLIC,CDWR:

Value1 = 1

Value2

Number of cyclic sectors

Value3

Number of solutions in cyclic space

Value4

Harmonic index of this load

Value5

Cyclic coordinate system

Value6

- < 0 or Static: only solve for given harmonic indices from CYCOPT, HIND
- > 0: tolerance for the Fourier load

Value1 = 2

Value2

Cyclic edge type (0 = undefined; 1 = areas; 10 = lines; 100 = keypoints; 1000 = nodes)

Value3

0 or blank

Value4

Maximum possible harmonic index

Value5

Force load coordinate system (1 = global coordinate system; 0 = cyclic coordinate system)

Value6

Inertia load coordinate system (1 = global coordinate system; 0 = cyclic coordinate system)

Value1 = 3 - 22

Value2 - Value6

Cyclic edge constraint equation/coupling degree of freedom (0 = all available degrees of freedom; otherwise bitmap) for pair IDs 1-5

(Repeat as necessary for other pair IDs (Value1 = 4 - 22))

```
Value1 = 23 - 30
```

Cyclic harmonic index bit bins (each bin holds 32 harmonic indices by 5 containers corresponding to Value2 - Value6)

Value2 - Value6

Cyclic harmonic index bits (0 = solve for harmonic index; nonzero values indicate skipped harmonic indices)

Value1 = 31

Value2

Max node number in base sector

Value3

Max element number in base sector

Value4

Number of defined nodes in base sector

Value5

Number of defined elements in base sector

Value1 = 32

Value2

/CYCEXPAND number of sectors to expand (total)

Value3

Number of edge component pairs

Value4 - Value8

/CYCEXPAND number of sectors to expand (per window)

Value1 = 33

Not used

Value1 = 34

Cyclic **CSYS** coordinate system integer data (part 1)

Value2

Theta singularity key

Phi singularity key

Value4

Coordinate system type

Value1 = 35

Cyclic CSYS coordinate system integer data (part 2)

Value2

Coordinate system number

Value3

Not used (defaults to 0)

Value4

Not used (defaults to 0)

Value1 = 36

Value2

Number of user-defined cyclic edge pair components

Value3

Rotate cyclic edge nodes into cyclic coordinate system (0 = rotate edge nodes (default); 1 = do not rotate edge nodes)

Value4

NLGEOM flag (0 = no **NLGEOM** effects (default); 1 = include **NLGEOM** effects)

Value5

Sector edge display key $(-1 = \text{suppresses display of edges between sectors even if the cyclic count varies between active windows; <math>0 = \text{averages stresses or strains across sector boundaries}$. This value is the default (although the default reverts to 1 or ON if the cyclic count varies between active windows); 1 = no averaging of stresses or strains occurs and sector boundaries are shown on the plot)

Value1 = 101

Value2

Sector angle (degrees)

XYZ tolerance input for matching low/high nodes

Value4

Angle tolerance for matching low/high nodes (degrees)

Value5

Tolerance in the element coordinate system for unequal meshes

Value1 = 102 - 104

Cyclic **CSYS** coordinate system double precision data (part 1)

Value2 - Value4

Coordinate system transformation matrix (total of 9 values)

Value1 = 105

Cyclic **CSYS** coordinate system double precision data (part 2)

Value2 - Value4

Origin location (XYZ)

Value1 = 106

Cyclic **CSYS** coordinate system double precision data (part 3)

Value2

Used for elliptical, spheroidal, or toroidal systems. If CSYS = 1 or 2, Value2 is the ratio of the ellipse Y-axis radius to X-axis radius (defaults to 1.0 (circle))

Value3

Used for spheroidal systems. If CSYS = 2, Value3 is the ratio of ellipse Z-axis radius to X-axis radius (defaults to 1.0 (circle))

Value1 = 107

Cyclic **CSYS** coordinate system double precision data (part 4)

Value2

First rotation about local Z (positive X toward Y)

Value3

Second rotation about local X (positive Y toward Z)

Third rotation about local Y (positive Z toward X)

Value1 = 121

Value2

Root of component names defining low and high ranges

Value1 = 122

Value2

Cyclic low/high xref array parameter name (node)

Value1 = 123

Value2

Cyclic low/high xref array parameter name (line)

Value1 = 124

Value2

Cyclic low/high xref array parameter name (area)

Value1 = 125

Value2

The component name of the elements to expand (see /CYCEXPAND,,WHAT)

Value1 = 201

Value2

Total number of modes extracted during a cyclic modal solve. This value is only available after call to **CYCCALC**.

Value3

Mode superposition flag to limit results written to .MODE and .RST files

Value4

Excitation engine order

Value1 = 202

Value2

Type of mistuning (1 = stiffness; 2 = mass; 3 = both; -1 = use user macro CYCMSUPUSERSOLVE)

Cyclic mode superposition restart flag (1 = new frequency sweep; 2 = new mistuning parameters; -1 = form blade superelement and stop)

Value4

Cyclic mode superposition key to perform complex modal analysis of reduced system

Value5

Number of CMS modes for mistuned reduced order model (see CYCFREQ,BLADE)

Value1 = 203

Value2

Array name for aerodynamic coupling coefficients

Value1 = 204

Unused

Value1 = 205

Value2

The name of the nodal component containing the blade boundary nodes at the blade-to-disk interface (see **CYCFREQ**,BLADE). This is used for cyclic mode superposition analyses that include mistuning or aero coupling.

Value1 = 206

Value2

The name of the element component containing the blade superelements (see **CYC-FREQ**,BLADE). This is used for cyclic mode superposition analyses that include mistuning or aero coupling.

Value1 = 207

Value2

The name of the array holding stiffness mistuning parameters

Value1 = 208

Unused

Value1 = 209

Rotational velocity from the base linear perturbation analysis.

X-component of rotational velocity

Value3

Y-component of rotational velocity

Value4

Z-component of rotational velocity

Value1 = 210

Value2

Beginning of frequency range for CMS modes (see **CYCFREQ**,BLADE). This is used for cyclic mode superposition analyses that include mistuning or aero coupling.

Value3

End of frequency range for CMS modes (see **CYCFREQ**,BLADE). This is used for cyclic mode superposition analyses that include mistuning or aero coupling.

Value1 = 211

Value2

Number of modes for a cyclic mode superposition damped modal solve

Value3

Beginning of frequency range for cyclic mode superposition damped modal solve

Value4

End of frequency range for cyclic mode superposition damped modal solve

3.2.7. EBLOCK Command

EBLOCK defines a block of elements. The command syntax is:

```
EBLOCK, NUM_NODES, Solkey Format
```

NUM_NODES

The number of nodes to be read in the first line of an element definition.

Solkey

The solid model key. The element is part of a solid model if the keyword SOLID appears here. When Solkey = SOLID, Field 8 (the element shape flag) may be left at zero, and Field 9 is the number of nodes defining this element.

Format

Data descriptors defining the format.

The format of the element block is as follows for the SOLID format:

- Field 1 The material number.
- Field 2 The element type number.
- Field 3 The real constant number.
- Field 4 The section ID attribute (beam section) number.
- Field 5 The element coordinate system number.
- Field 6 The birth/death flag.
- Field 7 The solid model reference number.
- Field 8 The element shape flag.
- Field 9 The number of nodes defining this element if Solkey = SOLID; otherwise, Field 9 = 0.
- Field 10 Not used.
- · Field 11 The element number.
- Fields 12-19 The node numbers. The next line will have the additional node numbers if there are more than eight.

The format without the SOLID keyword is:

- Field 1 The element number.
- Field 2 The type of section ID.
- Field 3 The real constant number.
- Field 4 The material number.
- Field 5 The element coordinate system number.
- Fields 6-15 The node numbers. The next line will have the additional node numbers if there are more than ten.

The final line of the block will be a -1 in field 1.

If you are in the GUI, the **EBLOCK** command must be contained in an externally prepared file and read into Mechanical APDL (via **CDREAD**, /INPUT, or other commands).

The **EBLOCK** command is not valid in a ***DO** loop.

3.2.8. EN Command

EN is used to define an element . If an element contains more than eight nodes, the **EN** command is repeated until all nodes are defined. The command format in Jobname . CDB is:

 $\verb|EN,UNBL|, Type|, NUMN|, I1, I2, I3, I4, I5, I6, I7, I8|\\$

Type

The type of data to be defined. Valid labels are ATTR (read in element attributes), and NODE (read in nodes defining the element).

NUMN

The number of nodes.

11,12,13,1415,16,17,18

The integer values to be read:

- If Type is ATTR, the integer values are the element attributes. Attributes are in the order: NUMN,MAT,TYPE,REAL,SECNUM,ESYS,NUMELEM,SOLID,DEATH,EXCLUDE
- If Type is NODE, the integer values are the node numbers.

3.2.9. LOCAL Command

LOCAL defines a local coordinate system. The command format in Jobname. CDB is:

LOCAL, UNBL, Type, NCSY, CSYTYP, VAL1, VAL2, VAL3

Туре

The type of data to be defined. Valid labels are LOC (read in system origin), ANG (read in rotation angles), and PRM (read in system parameters).

NCSY

The coordinate system reference number.

CSYTYP

The coordinate system type (0, 1, 2, or 3).

VAL1,VAL2,VAL3

Values to be read:

- If Type is LOC, values are the system origin in global Cartesian coordinates.
- If Type is ANG, values are the rotation angles in degrees.
- If Type is PRM, values are the first and second parameters of the system.

3.2.10. M Command

M defines a master degree of freedom. The command format in Johname . CDB is:

M, UNBL, NODE, Dlab

NODE

The node number

Dlab

The degree-of-freedom label

3.2.11. MPDATA Command

MPDATA defines a material property data table. You repeat the command until all properties are defined. The command format in Jobname . CDB is:

MPDATA, UNBL, LENGTH, Lab, MAT, STLOC, VAL1, VAL2, VAL3

LENGTH

The total number of temperatures in the table.

Lab

The material property label. (For valid labels, see MP.)

MAT

The material reference number.

STLOC

The starting location in the table for the next three property values.

VAL1,VAL2,VAL3

Property values assigned to three locations in the table starting at STLOC.

3.2.12. MPTEMP Command

MPTEMP defines a temperature table. You repeat the command until all temperature values are defined. The command format in Jobname . CDB is:

MPTEMP, UNBL, LENGTH, STLOC, TEMP1, TEMP2, TEMP3

LENGTH

The total number of temperatures in the table

STLOC

The starting location in the table for the next three temperature values

TEMP1,TEMP2,TEMP3

Temperatures assigned to three locations in the table starting at STLOC

3.2.13. N Command

If the UNBLOCKED option is used with **CDWRITE**, then **N** defines a node. The command format in Jobname. CDB is:

N, UNBL, Type, NODE, SOLID, PARM, VAL1, VAL2, VAL3

Type

The type of data to be defined. Valid labels are LOC (read in coordinates) and ANG (read in rotation angles).

NODE

The node number.

SOLID

The solid model reference key. Not present for Type= ANG.

PARM

Line parameter value (0 if not on line). Not present for Type= ANG.

VAL1,VAL2,VAL3

Values to be read:

- If Type is LOC, values are the coordinates in the global Cartesian system.
- If Type is ANG, values are the rotation angles in degrees.

3.2.14. NBLOCK Command

NBLOCK defines a block of nodes. This is the recommended method for inputting nodes into the Mechanical APDL database. The command syntax is:

NBLOCK, NUMFIELD, Solkey, NDMAX, NDSEL Format

NUMFIELD

The number of fields in the blocked format.

Solkey

The solid model key. The node is part of a solid model if the keyword SOLID appears here.

NDMAX

The maximum node defined.

NDSEL

The number of nodes written.

Format

Data descriptors defining the format.

The format of the node block is as follows:

- Field 1 Node number.
- Field 2 The solid model entity (if any) in which the node exists (if SOLID key).
- Field 3 The line location (if the node exists on a line and if SOLID key).
- Field 4 6 The nodal coordinates.
- Field 7 9 The rotation angles (if NUMFIELD > 3).

Only the last nonzero coordinate/rotation is output; any trailing zero values are left blank.

The final line of the block is always an **N** command using a -1 for the node number.

The following example shows a typical **NBLOCK** formatted set of node information. Note that this example has no rotational data. It contains only the first six fields.

```
NBLOCK, 6, SOLID,
                      849,
                                 723
(3i9,6e21.13e3)
                 0
                          0 8.7423930292124E-001 7.1843141243360E-001 8.2435547360131E-001
        1
        3
                 0
                          0 9.2314873336026E-001 9.3459943382943E-001 4.8406643591666E-001
                 0
        4
                          0 1.1410427242574E+000 7.6883495387624E-001 2.5867801436812E-001
      847
                 0
                          0 6.2146469267794E-001 8.0122597436764E-001 8.1352232529497E-001
      848
                 0
                          0 8.1179373384170E-001 6.6711479947438E-001 7.6547291135454E-001
                          0 7.4952223718564E-001 7.6089019544242E-001 7.4112247735703E-001
      849
                 0
N, UNBL, LOC,
```

If you are in the GUI, the **NBLOCK** command must be contained in an externally prepared file and read into Mechanical APDL (**CDREAD**, /**INPUT**, or other commands).

The **NBLOCK** command is not valid in a ***DO** loop.

3.2.15. *PREAD Command

```
*PREAD,Par,NUMVALS
Format
END PREAD
```

Par

Alphanumeric name to identify this parameter.

NUMVALS

Number of values.

Format

Data descriptor defining the format of the subsequent lines (as needed). The format is always (4g20.13).

3.2.16. R Command

R defines a real constant set. You repeat the command until all real constants for this set are defined. The command format in Jobname . CDB is:

R, UNBL, NSET, Type, STLOC, VAL1, VAL2, VAL3

NSET

The real constant set reference number.

Type

The type of data to be defined. LOC is the valid label.

STLOC

The starting location in the table for the next three constants.

VAL1,VAL2, VAL3

Real constant values assigned to three locations in the table starting at STLOC.

3.2.17. RLBLOCK Command

RLBLOCK defines a real constant set. The real constant sets follow each set, starting with Format1 and followed by one or more Format2's, as needed. The command format is:

RLBLOCK, NUMSETS, MAXSET, MAXITEMS, NPERLINE Format1
Format2

NUMSETS

The number of real constant sets defined

MAXSET

Maximum real constant set number

MAXITEMS

Maximum number of reals in any one set

NPERLINE

Number of reals defined on a line

Format1

Data descriptor defining the format of the first line. For **RLBLOCK**, this value is always (2i8,6g16.9). The first i8 is the set number, the second i8 is the number of values in this set, followed by up to six real constant values.

Format2

Data descriptors defining the format of the subsequent lines (as needed); this is always (7g16.9).

The real constant sets follow, with each set starting with Format1, and followed by one or more Format2's as needed.

RLBLOCK is not valid in a ***DO** loop.

3.2.18. SE Command

SE defines a superelement. The command format in Jobname. CDB is:

SE, UNBL, File, , , TOLER, TYPE, ELEM

File

The name (case-sensitive) of the file containing the original superelement matrix created by the generation pass (Sename . SUB).

TOLER

Tolerance for determining whether use-pass nodes are noncoincident with master nodes having the same node numbers. Default = 0.0001.

TYPE

Element type number.

ELEM

Element number.

This command command also appears in the *Command Reference*. The format shown here contains information specific to the **CDREAD/CDWRITE** file.

3.2.19. SECBLOCK Command

SECBLOCK for Beams

SECBLOCK retrieves all mesh data for a user-defined beam section as a block of data. You repeat the command for each beam section that you want to read. The command format is:

```
SECBLOCK
Format1
Format2
Format3
```

Format1

The First Line section. The first value is the number of nodes, and the second is the number of cells.

Format2

The Cells Section. The first 9 values are the cell connectivity nodes. The 10th (last) value is the material ID (MAT).

Format3

The Nodes Section. This section contains as many lines as there are nodes. In this example, there are 27 nodes, so a total of 27 lines would appear in this section. Each node line contains the node's boundary flag, the Y coordinate of the node, and the Z coordinate of the node. Currently, all node boundary flags appear as 0s in a cell mesh file. Because all node boundary flags are 0, **SECBLOCK** ignores them when it reads a cell mesh file.

Sample User Section Cell Mesh File

Following is a sample excerpt from a custom section mesh file for a section with 27 nodes, 4 cells, and 9 nodes per cell:

First Line:	27	4			
Cells Section:		1 3 11 7 9 23 9 11 25 11 13 27	9 2 6 21 8 16 23 10 18 25 12 20	24 16 17	2 1 1 1
Nodes Section:	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	0.0 0.025 0.05 5.0175 19.98 20.00	0.0 0.0 0.0 0.0 10.00 10.00		

SECBLOCK for Shells

SECBLOCK can retrieve data for shell sections. The command format is:

```
SECBLOCK, N
TKn, MATn, THETAn, NUMPTn
```

N

Total number of layers. The second line (TKn, MATn, THETAn, NUMPTn) is repeated N times (once for each layer).

TKn

Layer thickness for layer number n

MATn

Material ID for layer number n (defaults to element material ID)

THETAn

Layer orientation angle for layer number n

NUMPTn

Number of integration points in layer number n

SECBLOCK is not valid in a ***DO** loop.

3.2.20. SFBEAM Command

SFBEAM defines a surface load on selected beam elements. Remaining values associated with this specification are on a new input line with a 4(1pg16.9) format. The command format in Jobname . CDB is:

SFBEAM, ELEM, LKEY, Lab, UNBL, DIOFFST, DJOFFST

ELEM

The element number.

LKEY

The load key associated with these surface loads.

Lab

A label indicating the type of surface load. PRES (for pressure) is the only valid label.

DIOFFST

Offset distance from node I.

DJOFFST

Offset distance from node J.

3.2.21. SFE Command

If the UNBLOCKED option is used with **CDWRITE**, then **SFE** defines a surface load. Values associated with this specification are on a new input line with a 4(1pg16.9) format. The command format in Jobname. CDB is:

SFE, ELEM, LKEY, Lab, KEY, UNBL

ELEM

The element number.

LKEY

The load key associated with this surface load.

Lab

A label indicating the type of surface load. (For a list of load labels, see SFE.)

KEY

A value key. The possible values and meaning of each value depend on the specified load label. (See the KVAL argument of **SFE** for a list of value keys based on load label.)

3.2.22. SFEBLOCK Command

SFEBLOCK defines a block of element surface loads. This is the recommended method for inputting element surface loads into the Mechanical APDL database. The command syntax is:

SFEBLOCK, NUMFIELD, Lab, ELMAX, ELSEL, TAB Format

NUMFIELD

The number of fields in the blocked format.

Lab

The surface load label used to describe the block data. (For a list of load lables, see SFE.)

ELMAX

The maximum element number defined.

ELSEL

The number of selected elements.

TAB

Key for tabular input:

0 – Non-tabular input.

1 - Tabular input.

Format

Data descriptors defining the format.

The format of the surface load block is as follows:

- Field 1 Element number.
- Fields 2 Load key or face number associated with the surface load.
- Fields 3 Value key. (See the KVAL argument of SFE for a list of value keys based on load label.)
- Fields 4-7 Surface load values to apply to each element.

The final line of the block format is always an SFE command with -1 for the element number.

The following example shows a typical **SFEBLOCK** formatted set of body-force loads (using non-tabular input) that define a convection load (CONV) characterized by film coefficient and bulk temperature.

```
SFEBLOCK, 4, CONV,
                                   5,0
(i9,i4,i4,6(pg16.9))
       3 1 1 10.0000000
                                   10.0000000
                                                   0.00000000
                                                                  0.00000000
       3
           1
               2 300.000000
                                   300.000000
                                                  0.00000000
                                                                  0.00000000
        4
           1
               1
                  6.50000000
                                   6.50000000
                                                  0.00000000
                                                                  0.00000000
           1
               2 146.153800
                                                  0.00000000
                                                                  0.00000000
                                   146.153800
          1
              1 3.50000000
                                   3.50000000
                                                   0.00000000
                                                                  0.00000000
       5
           1
               2 300.000000
                                   300.000000
                                                   0.00000000
                                                                  0.00000000
SFE, end, LOC,
                   -1,
```

In the GUI, **SFEBLOCK** must be contained in an externally prepared file and read into Mechanical APDL (via **CDREAD**, /INPUT, or other commands).

SFEBLOCK is not valid in a ***DO** loop.

Part 2: Guide	to User-Pro	grammable	Features

Chapter 1: Understanding User Programmable Features (UPFs)

The Mechanical APDL program has an open architecture, allowing you to write your own routines or subroutines in C, C++, or FORTRAN and either link them to the program or use them as external commands. Some standard program features originated as user customizations, also known as user programmable features (UPFs).

You can take advantage of user customization if you are licensed for the ANSYS Mechanical Enterprise family of products (ANSYS Mechanical Enterprise, ANSYS Mechanical Enterprise PrepPost, and ANSYS Mechanical Enterprise Solver).

Other ANSYS, Inc. products do not support customization of the Mechanical APDL program.

For more information about compilers, see the ANSYS, Inc. installation guide specific to your operating system:

ANSYS, Inc. Linux Installation Guide ANSYS, Inc. Windows Installation Guide

The following customization topics are available:

- 1.1. What Are User Programmable Features?
- 1.2. What You Should Know Before Using UPFs
- 1.3. Planning Your UPFs
- 1.4. Studying the Mechanical APDL User Routines
- 1.5. Programming in Languages Other Than FORTRAN
- 1.6. Developing UPFs: a Suggested Strategy
- 1.7. Include Decks
- 1.8. Choosing a Linking Method
- 1.9. Compiling and Linking UPFs on Linux Systems
- 1.10. Sharing Data Between User Routines
- 1.11. Compiling and Linking UPFs on Windows Systems
- 1.12. Activating UPFs
- 1.13. Running Your Custom Executable
- 1.14. Verifying Your Routines
- 1.15. Debugging Commands
- 1.16. Other Useful Commands
- 1.17. Generating Output
- 1.18. Reading Large Data Files More Rapidly

1.1. What Are User Programmable Features?

User programmable features are tools you can use to write your own routines. Using UPFs, you can tailor the Mechanical APDL program to your organization's needs. For instance, you may need to define a new material behavior, a special element, a contact interfacial model, or a modified failure criterion for composites.

UPFs provide the following capabilities:

- To read information into or retrieve information from the Mechanical APDL database, you can create subroutines and either link them into the program or use them in the external command feature (see Appendix A: Creating External Commands in Linux (p. 363) for more information about external commands). If you link these subroutines into Mechanical APDL, you are limited to 10 database access commands. Such commands, created through either method, operate at all levels of program operation, including the begin, preprocessor, general postprocessor, time-history postprocessor, and solution levels. For more information about accessing the database, see Accessing the Mechanical APDL Database (p. 283).
- Some UPF subroutines enable you to specify various types of loads, including BF or BFE loads, pressures, convections, heat fluxes, and charge densities. These routines are described under Subroutines for Customizing Loads (p. 246).
- Some UPF subroutines enable you to modify and monitor existing elements. For details, see Subroutines for Modifying and Monitoring Existing Elements (p. 192).
- Some UPF subroutines enable you to define the following material properties: plasticity, creep, swelling law, viscoplasticity, hyperelasticity, and layered element failure criteria. To see inputs and outputs for these routines, see Subroutines for Customizing Material Behavior (p. 198).
- For analyses involving contact, another set of UPF subroutines enables you to define contact properties, friction models, and interaction behaviors. To see inputs and outputs for these routines, see Subroutines for Customizing Contact Interfacial Behavior (p. 231).
- Several sets of UPFs enable you to define new elements and to adjust the nodal orientation matrix. See Creating a New Element (p. 149) for more information.
- You can call Mechanical APDL as a subroutine in a program you have written. To learn how, see Running Mechanical APDL as a Subroutine (p. 260).

1.2. What You Should Know Before Using UPFs

Before you do anything with linked UPFs, contact your on-site system support person to get the permissions needed to access the appropriate ANSYS, Inc. files.

The UPF subroutines are written in FORTRAN; some extensions are used. They contain comments intended to give you enough detail to develop your own versions of the subroutines.

User routines that can be modified have the term "USERDISTRIB" in the first line of the routine. These routines are provided with the ANSYS, Inc. distribution media. You can modify these routines to tailor the program to your specific needs. Certain other routines described in this document are not provided on the distribution media, but can be called using the provided header information.

To use UPFs successfully, you need strong working knowledge of the following:

- The Mechanical APDL program.
- The UPF subroutines themselves. Study the UPF subroutines before customizing them, and make sure that you fully understand the subroutines, as well as any applicable functions. Unless you review them carefully, a few UPF subroutines may seem like a maze with many logic paths to consider. You may have to set special variables correctly in order to run your customized Mechanical APDL solution without errors. Even if you have in-depth knowledge of the Mechanical APDL input and your desired outputs, you still need to ensure that everything that must be done in the UPF subroutines is done properly in your custom version.
- FORTRAN. Besides knowing how to write FORTRAN subroutines, you must be sure that the level of
 the FORTRAN compiler is as least as high as the level mentioned in your ANSYS, Inc. installation
 manual. For more information on FORTRAN compilers, please refer to the installation guide specific
 to your operating system (ANSYS, Inc. Linux Installation Guide or ANSYS, Inc. Windows Installation
 Guide). You also need to know what to do should the computer abort the program due to an arithmetic error, a file read error, a memory access error, and so on.
- The mathematics of the phenomenon you are planning to include.

Important

- UPFs are not available or will behave unpredictably in certain data center environments or on some
 hardware configurations. You should take special care when using UPFs on parallel systems. It is a
 good practice to verify your coding with single processing by using the -np,1 option before you run
 your analysis. For additional information, consult your installation guide or your on-site ANSYS systemsupport person
- Carefully consider whether you wish to use UPFs, especially if you are linking them into Mechanical
 APDL (rather than into a shared library for use as external commands). When you add your own
 routines to Mechanical APDL by either method, you are creating a customized, site-dependent version
 of the program. ANSYS, Inc. considers the use of UPFs a nonstandard use of the program, one that
 the ANSYS, Inc. Quality Assurance verification testing program does not cover. Therefore, you are
 responsible for verifying that the results produced are accurate and that your customizations do not
 adversely affect unchanged areas of the Mechanical APDL program.
- Although the flexibility that UPFs offer can be highly attractive, UPF usage is a complicated process
 that can introduce errors. Consider what you want your customizations to accomplish. You may be
 able to customize Mechanical APDL more easily and safely with macros than with UPFs.
- When using shared memory, all user-programmable features are supported except for common block variables in FORTRAN and external variables in C or C++. You should avoid overwriting the values of these variables by multiple cores at the same time.
- For Distributed ANSYS, all user-programmable features are supported except for global (often in common block) variables in FORTRAN and global (often external) variables in C or C++. You should avoid overwriting the values of these variables; they should have the same values across all cores used in the distributed solution.

For other guidelines for nonstandard uses of the Mechanical APDL program, see User-Programmable Features and Nonstandard Uses in the *Advanced Analysis Guide*.

1.3. Planning Your UPFs

UPFs can range from a simple element output routine for customized output to a complex user optimization. Before you start programming, ask yourself these questions:

- Does the capability you want already exist in Mechanical APDL? Remember, a capability may not be obvious at first, especially to a novice user.
- Does your proposed subroutine fit into the Mechanical APDL program architecture and specifications? For example, you can not program a user element that has more than 32 degrees of freedom per node.

Use your experience and judgment to answer these questions. If you need help to do so, consult your ANSYS Support Distributor. If you can respond "no" to the first question and "yes" to the second question, then the user routine you are planning will be both useful and feasible.

1.4. Studying the Mechanical APDL User Routines

Your ANSYS, Inc. distribution medium contains the source codes for all user routines:

- If you are running Mechanical APDL under Linux, the source code for the UPF routines resides in directory /ansys_inc/v211/ansys/customize/user/.
- If you are running Mechanical APDL under Windows, the source code for the UPF routines resides in directory Program Files\ANSYS Inc\V211\ansys\customize\user\.

Most of the user routines have at least simple functionality, so print out the routines of interest before you start programming. All source routines are concatenated onto file user.for.Delete the routines you do not want and make appropriate changes to the others.

1.5. Programming in Languages Other Than FORTRAN

If you wish to run Mechanical APDL with user customizations, the preferred method is to design and program your custom routine in FORTRAN. Although you can use languages other than FORTRAN, in each case FORTRAN must provide the interface to the rest of the Mechanical APDL program. If you do use a language other than FORTRAN, such as the C or C++, your code may require a FORTRAN shell.

You need to take care when calling FORTRAN subroutines from C or C++ subroutines. You must use the symbol associated with the FORTRAN subroutine when invoking the subroutine from a C or C++ function. This symbol typically differs slightly from the FORTRAN subroutine name, and is extremely system dependent.

On many Linux systems, you build this symbol name by taking the FORTRAN subroutine name, converting it to lower case, and appending an underscore. For example, the symbol name for the FORTRAN subroutine **HeapInquire** would be **heapinquire**. You would have to use the symbol **heapinquire** in the invoking C function to avoid an unsatisfied external reference when the program is linked.

Keep in mind that the instance described above is just an example. Compilers from different vendors may construct the symbols differently. Please consult the manuals for your specific compiler for information on how to call FORTRAN subroutines from C or C++ functions.

For more information on FORTRAN compilers please refer to the installation guide specific to your operating system (ANSYS, Inc. Linux Installation Guide or ANSYS, Inc. Windows Installation Guide).

1.6. Developing UPFs: a Suggested Strategy

When developing customizations for Mechanical APDL, you can avoid problems and reduce debugging time by following a gradual, orderly process. Start with a trivial test. Then, add a few changes at a time so that if something goes wrong, the error that caused the problem should be isolated and relatively easy to locate.

The example procedure below illustrates this type of gradual process. The example assumes that you are creating a new element using the method described in Creating a New Element by Directly Accessing the Program Database (p. 169). You develop and test it by performing these steps:

- 1. Get the applicable element subroutines for uel101 from the product distribution medium. Add a small change (such as a misspelling in an output heading), then compile and link the subroutines.
- 2. Using a production version of the program, run several analysis problems using various elements to form a base for comparison.
- 3. Run the same problem using USER101 on your custom version of the program.
- 4. Compare the results from Steps 2 and 3. If they show discrepancies other than the misspelled output heading, resolve them before you go on to Step 5.
- 5. Choose the standard Mechanical APDL element that most closely resembles your new custom element, and run some problems on a production version of Mechanical APDL using that element.
- 6. Modify the element subroutines to match the element you chose in Step 5. Then, compile and link those subroutines into a custom version of Mechanical APDL.
- 7. Again, compare the results from Steps 5 and 6. If they don't match, resolve the discrepancies before moving on to Step 8.
- 8. Modify your element subroutines to include the features you want. Then, compile and link the subroutines into a custom version of Mechanical APDL.
- 9. Test the changes with a series of increasingly complex problems for which you already know the answers.

1.7. Include Decks

In addition to the subroutines and functions described in this chapter, most of the include decks (files with the extension .inc) used by Mechanical APDL are on your product distribution medium. The include decks, also called *commons*, contain important but relatively small amounts of data. The program also

handles large amounts of data using various access routines (GET and PUT), described elsewhere in this document.

Note:

When you compile a user-programmable feature (UPF) in a shared library (ANSUSER-SHARED.BAT), you cannot access the common block variables. (All such variables will return a value of zero.)

To insert include decks in a subroutine or function, use the INCLUDE (or an analogous) statement. *Do not modify an include deck under any circumstances*. The following table lists some of the more commonly used include files and the definitions they contain:

Include File	Description
acel- cm.inc	Contains accelerations and angular velocities
ansys- def.inc	Defines general Mechanical APDL parameters. You must include this common to retrieve the parameter values of MEM_INTEGER, MEM_DOUBLE, MEM_COMPLEX, or MEM_REAL.
cmopt.inc	Contains optimization variables
ech- prm.inc	Defines parameters for element characteristics
el- ccmt.inc	Defines element characteristics (comments only)
ele- com.inc	Contains element-specific information
el- parm.inc	Defines pointers for the element data array
eluc- om.inc	Defines the element degree of freedom pointers
ety- com.inc	Element type data
imp- com.inc	Used by all routines and functions in the program
outp- cm.inc	Defines output control information
soptcm.inc	Contains solution options and keys
stack.inc	Defines stack storage. You must include this common in any routines that access stack space.
step- cm.inc	Contains load step information
us- vrcm.inc	Defines storage of user-defined variables

1.8. Choosing a Linking Method

After you make your changes to the user routines supplied on your product distribution medium, you can either:

- Link your routines into shared libraries (as discussed starting in Appendix A: Creating External Commands in Linux (p. 363)), or
- Compile and link your custom routines into the Mechanical APDL program itself. This is discussed for Linux systems in Compiling and Linking UPFs on Linux Systems (p. 133) and for Windows systems in Compiling and Linking UPFs on Windows Systems (p. 138). You may need superuser or root privileges to run the procedure that does the linking.

For both Windows and Linux platforms, three methods of compiling and linking are available:

- /UPF command
- ANSUSERSHARED script (creates a shared library on Linux or a dynamic-link library on Windows)
- ANS_ADMIN211 Utility

The /UPF command method is typically used by individuals wanting to occasionally use their customized code for certain runs. The advantages of this method are that it is very easy to use and the source code can be displayed in the output file.

The shared library (Linux) and dynamic link library (Windows) methods are typically used to run Mechanical APDL with frequently used user-libraries or third-party libraries (material libraries, and so on). This method is advantageous if customized code is frequently used or shared with other users.

The **ANS_ADMIN211** method is useful for someone wanting to create a permanently changed Mechanical APDL executable which will be used by many users, or used most of the time. Companies that validate their user-customized code might want to consider this option.

In some cases, you might want to combine two of the methods of compiling and linking. The program allows you to combine the **ANS_ADMIN211** method with either the /**UPF** command method or the ANSUSERSHARED method. Note that the /**UPF** command method cannot be combined with the ANSUSERSHARED method.

As an example of combining these methods, you might first create a custom executable with **ANS_ADMIN211** that contains user creep laws. Then, you might use the **ANS_USER_PATH** environment variable to include a user material (or third-party library) created with the ANSUSERSHARED method.

For detailed compiling and linking procedures, see Compiling and Linking UPFs on Linux Systems (p. 133) and Compiling and Linking UPFs on Windows Systems (p. 138).

1.9. Compiling and Linking UPFs on Linux Systems

There are three methods that you can use to link your custom routines into Mechanical APDL:

- Using the /UPF Command (p. 134)
- Creating a Shared Library (p. 135)

• Using the ANS_ADMIN Utility (p. 135)

The source files for the user routines reside in the following subdirectory: /ansys_inc/v211/ansys/customize/user/

For all three methods, you can write your user routines in one language or a combination of languages (FORTRAN, C, and C++). Note that when using a combination of languages, you are responsible for the calling interfaces between languages (see Programming in Languages Other Than FORTRAN (p. 130)).

The ANSYS, Inc. Linux Installation Guide lists the compilers that are required for specific platforms.

Note:

You will need all the compilers specified in the Installation Guide specific to your operating system to use these user programmable features, including GNU GCC 8.2.0. This specific compiler can be downloaded from the ANSYS, Inc. customer site. See Downloading and Installing the GCC Compiler (p. 136) for details.

1.9.1. Using the /UPF Command

The /UPF command offers the simplest method for linking user programmable features into Mechanical APDL. The format of the command is:

```
/UPF, RoutineName
```

where <code>RoutineName</code> is the name of a user routine (filename.ext) that you want to link. The specified routine must reside in the current working directory.

To use this method start Mechanical APDL in batch mode and include one or more /UPF commands in the specified input listing. When the program reads the input and detects /UPF, Mechanical APDL will be relinked automatically.

You can include /UPF anywhere in your input file, and you can repeat /UPF as many times as needed to include multiple user routines in the relinked version. Any user routine can be linked using this method.

When you run a user-linked version of the program by this method, the output includes the following:

```
NOTE - This Mechanical APDL version was linked by /UPF with n user supplied routine(s).
```

where n indicates the number of routines specified by **/UPF** commands. The routines that have been linked will be included in the output listing.

Example Using Mixed Languages

The following directory contains an example of using the /UPF command method to link user routines that are written in mixed languages (FORTRAN, C, C++):

```
/ansys_inc/v211/ansys/custom/user/<platform>/Examples
```

The README.txt file in this directory contains complete instructions on running this example. This is a simple, automated test that verifies whether compilers are correctly installed and configured.

1.9.2. Creating a Shared Library

You can also set up UPFs on some Linux systems through a shared library. All FORTRAN files (files ending with .F), C files (files ending with .C), and C++ files (files ending in .Cpp) that you want to include in your shared library should reside in your working directory. To compile all * .F, * .C, and * .Cpp routines, issue the following command:

```
/ansys_inc/v211/ansys/customize/user/ANSUSERSHARED
```

If the compile was successful, you will be asked if a shared file is to be created. Choose **Yes** to create a shared library named libansuser.so.

To use this library, set the ANS_USER_PATH environment variable to point to the working directory where the libansuser shared library resides. Use one of the following commands, depending on the Linux shell you are using:

```
setenv ANS_USER_PATH workingdirectory
```

or

export ANS_USER_PATH=workingdirectory

When you run a user-linked version of Mechanical APDL, the output echos the value of **ANS_USER_PATH** and will include the following:

```
NOTE: This Mechanical APDL version was linked by Licensee
```

To return to the original version of ANSYS, unset the **ANS USER PATH** environment variable.

You can use another environment variable, **ANS_USER_PATH_211**, to point to a shared library specific to Release 2021 R1. **ANS_USER_PATH_211** supersedes **ANS_USER_PATH**. This allows you to set up and use more than one shared library containing UPFs. To change back to the shared library specified by **ANS_USER_PATH**, simply unset the **ANS_USER_PATH_211** environment variable.

ANSYS, Inc. recommends using the ANSUSERSHARED script as a template to try compilers that are not supported by ANSYS, Inc., such as the GNU compilers. To do so, edit the ANSUSERSHARED script, making changes to the appropriate platform logic. Note that if you do use compilers other than those listed in the ANSYS Installation and Configuration Guide specific to your operating system, you will need to debug (that is, find missing libraries, unsatisfied externals, etc.) them yourself. ANSYS, Inc. does not provide assistance for customers using unsupported compilers or if the resulting objects are not compatible with the executable(s) as distributed.

1.9.3. Using the ANS_ADMIN Utility

As mentioned previously, the source files for the user routines reside in subdirectory /an-sys_inc/v211/ansys/customize/user/. If you modify any of these subroutines, you can select the **Relink ANSYS** option from **ANS_ADMIN211** utility to link these changes. This method creates a custom Mechanical APDL executable.

The **Relink ANSYS** option compiles all FORTRAN files (files ending with .F), C files (files ending with .c), and C++ files (files ending in .cpp) in the current working directory. The procedure then loads all object files (files ending with .o) along with the default Mechanical APDL objects and libraries in

/ansys_inc/v211/ansys/customize/user/<platform>. For Distributed ANSYS the location is: /ansys_inc/v211/ansys/customize/user/<platform>/dis/native, where <platform> is replaced by the folder representative of your operating system. The new executable file created will be named ansyscust.e and will reside in the working directory.

When you run a user-linked version of Mechanical APDL, the output includes the following:

```
NOTE: This Mechanical APDL version was linked by Licensee
```

If you intend to run the linked version of Mechanical APDL in a distributed environment (for example, on a cluster), the executable (ansyscust.e) must reside in the same directory path on all systems. However, you need to link it on only one system; you can then copy the executable to the other systems.

1.9.4. Downloading and Installing the GCC Compiler

In order to link UPFs on a Linux system, we recommend this version of the GCC compiler: GNU GCC 8.2.0. As a convenience, you can download this compiler from the ANSYS, Inc. customer site and extract the necessary files using these steps:

- From the ANSYS customer site, www.ansys.com/customercommunity, click **Downloads > Current** Release.
- 2. Select the **Linux x64** operating system.
- 3. Select **Primary Packages** from the **Select Download Type** drop-down menu.
- 4. Expand the **Tools** menu.
- 5. Click the **GCC Compiler** download option.
- 6. Select your desired download directory and click **Save**.

This will download the compressed zip file linux-toolchain-8.2.0-01.zip.

7. Create a linux-toolchain-8.2.0-01 folder to contain the contents of the zip file:

```
mkdir linux-toolchain-8.2.0-01
```

8. Extract the file within the location created in step 7.

```
unzip linux-toolchain-8.2.0-01.zip
```

9. Add the location to your PATH, or set **ANS_GCC_PATH** for access to the GCC 8.2.0 compiler. For example:

```
setenv PATH <installed_location>/linux-toolchain-8.2.0-01/bin:$PATH (for csh/tcsh shell)
export PATH=<installed_location>/linux-toolchain-8.2.0-01/bin:$PATH (for sh/bash shell)
```

```
or
```

```
\label{location} {\tt setenv ANS\_GCC\_PATH < installed\_location>/linux-toolchain-8.2.0-01/bin (for csh/tcsh shell)} \\ {\tt export ANS\_GCC\_PATH=< installed\_location>/linux-toolchain-8.2.0-01/bin (for sh/bash shell)} \\ {\tt occupation>/linux-toolchain-8.2.0-01/bin (for sh/bash shell)} \\ {\tt occupation>/linux-to
```

The GCC version should display gcc (ansys-20190911) 8.2.0.

1.10. Sharing Data Between User Routines

Mechanical APDL allows more than one user routine in a single run. You can issue multiple /**UPF** commands in the input file to activate the routines.

Using multiple routines simultaneously may require sharing data generated by one routine with another. The easiest method for doing so is to use common-block (or global) variables.

In Linux, a single shared library contains all compiled user routines and data sharing is straightforward. In Windows, however, each user routine is built into a separate dynamic link library (DLL); to share data, functions and data must be explicitly exported and imported.

The userdata (p. 259) subroutine enables you to store the common-block functionality and data. You can edit usercm (p. 260) (included in userdata) to add common-block variables. The compiler uses !DEC\$ ATTRIBUTES DLLEXPORT and !DEC\$ ATTRIBUTES DLLIMPORT to indicate which functions and common-block variables to export or import, respectively. Both commands are valid for the supported Intel FORTRAN and C compilers.

Example 1.1: Creating and Exporting Functions and Common Block Variables

```
c usercm.inc
*comdeck,usercm USERDISTRIB
!DEC$ ATTRIBUTES DLLEXPORT :: /usercm/
        common /usercm/ userdatsz,userdatptr
        pointer (userdatptr,userdataarray)
        integer userdatsz
  double precision userdataarray(*)
С
c userdat.F . Sample function exported for use in other subroutines
      function getusercmvals(iloc,sz,outdata)
!DEC$ ATTRIBUTES DLLEXPORT :: getusercmvals
#include "usercm.inc"
                      iloc,sz,getusercmvals,iX
      integer
      double precision outdata(*)
   double precision userdataloc(*)
      pointer (userdatptr,userdataloc)
      if (iloc.lt.1.or.iloc+sz-1.gt.userdatsz) then
         getusercmvals = 0
      else
    do iX=iloc,iloc+sz-1,1
   outdata(iX) = userdataloc(iX)
    enddo
          getusercmvals = 1
      endif
      return
      end
```

Example 1.2: Importing and Using Functions

Imported functions are added to the interface section. Common blocks are also imported as needed.

```
INTERFACE
FUNCTION getusercmvalsz ()
!DEC$ ATTRIBUTES DLLIMPORT :: getusercmvalsz
integer getusercmvalsz
END
return
end
END INTERFACE
```

```
!DEC$ ATTRIBUTES DLLIMPORT :: /usercm/
    common /usercm/ userdatsz,userdatptr
    pointer (userdatptr,userdataarray)
    integer userdatsz
    double precision userdataarray(*)
```

Using common-block variables in shared memory requires care. Multiple threads started by the executable access and share the same memory location and can overwrite each other's values. To minimize conflict, allocate enough space for each thread and avoid writing to the same location at the same time.

Example 1.3: Safely Allocating Separate Data for Each Thread

One double-precision location exists in the common block. Enough space is allocated for <code>numberof-processors*</code> 1 locations. Functions <code>pplock</code> and <code>ppunlock</code> are used when initializing the memory or values.

When setting the value, write to the iy memory location only (used for that specific thread). Even if other threads access the same common block, they do not modify that memory location.

```
c Initialize common-block values in user routine 1 such as USolBeg.F
  call pplock(LOCKUPF)
  inumprocs = ppinqr(PP_NPROCS)
  call initusercmvals(inumprocs)
     val1(:) = 0.0d0
 iy = setusercmvals(ix, inumprocs,val1)
  call ppunlock(LOCKUPF)
C
C
c Set common-block value to time in user routine 2
c Go to location allocated for the iy the thread
 iy = ppproc()+1
  val1 = time
c Set just the iy-th value.
 iy = setusercmvals(iy,isize,val1)
c Get saved value from common block in user routine 3
  iproc = ppproc()+1
  isize = 1
  call getusercmvals(iproc,isize,dFldTime)
```

For information about the pplock (p. 345), ppinqr (p. 344), ppunlock (p. 345), and ppproc (p. 345) subroutines used in Example 1.3: Safely Allocating Separate Data for Each Thread (p. 138), see Subroutines for Your Convenience (p. 343).

1.11. Compiling and Linking UPFs on Windows Systems

There are three methods that you can use to link your custom routines into the Mechanical APDL program:

```
Use the /UPF command (p. 140)
Create a dynamic-link library (p. 142)
```

Use the ANS_ADMIN Utility (p. 144)

The source files for the user routines reside in the following subdirectory: Program Files\ANSYS Inc\V211\ansys\customize\user\.

The user programmable features are loaded onto the system only if you perform a custom installation and choose to install the customization tools. If you intend to modify any of the user routines, make a duplicate copy of the Program Files\ANSYS Inc\V211\ansys\customize\user\ directory to preserve the original files for later use, if necessary.

For all three methods, you can write your user routines in one language or a combination of languages (FORTRAN, C, and C++). Note that when using a combination of languages, you are responsible for the calling interfaces between languages (see Programming in Languages Other Than FORTRAN (p. 130)).

The ANSYS, Inc. Windows Installation Guide lists the compilers that are required for Windows systems.

Note:

You will need all the compilers specified in the Installation Guide specific to your operating system to use these user programmable features. ANSYS, Inc. does not provide assistance if customers are using unsupported compilers, or if the resulting objects are not compatible with the Mechanical APDL executable(s) as distributed.

Before linking UPFs, make sure that the INCLUDE, LIB, and PATH environment variables are set correctly for the required C/C++ and Intel FORTRAN compilers.

Visual Studio 2017 Professional is also required for linking user programmable features on Windows platforms. In Visual Studio 2017, C++ is not installed by default. To install C++, you must select **Custom Install** and select **Common Tools for C++ for 2017**.

Before using any of the methods described below for linking UPFs, open the following Command Prompt window if you have Visual Studio 2017 Professional installed:

Start > All apps > Visual Studio 2017 > Developer Command Prompt for VS2017

Note:

For all three linking methods, you might have issues with write/modify access if you use the default working directory: Program Files\ANSYS Inc\V211\ansys\custom\user\. If the linking operation errors out or fails, try running the required applications as an administrator by right clicking and choosing "Run as administrator" while launching each application.

For all three linking methods, you can set the **ANS_USER_PATH** environment variable to specify the working directory where the created UPF DLLs reside. You can use another environment variable, **ANS_USER_PATH_211**, to point to a set of UPF DLLs specific to Release 2021 R1. **ANS_USER_PATH_211** supersedes **ANS_USER_PATH**. This allows you to set up and use more than one set of UPF DLLs. To change back to the location specified by **ANS_USER_PATH**, simply unset the **ANS_USER_PATH_211** environment variable.

1.11.1. Using the /UPF Command

The /UPF command offers the simplest method for linking user programmable features into Mechanical APDL. The format of the command is:

```
/UPF, RoutineName
```

where RoutineName is the name of the user routine (filename.ext) that you want to link. The specified routine must reside in the current working directory.

Following are prerequisites for using the /UPF command method on a Windows system:

- A script named findUPF.bat is used to detect the /UPF command. You must include the path to this script in your system PATH variable. This script is typically located in Program Files\Ansys Inc\V211\ansys\bin\<platform> where <platform> is a directory that uniquely identifies the hardware platform version: "Winx64" for 64-bit Windows.
- Before starting Mechanical APDL, you must set the ANS_USE_UPF environment variable to TRUE. This causes the program to search for /UPF in the input file. This environment variable is required only on Windows systems and only when using the /UPF command method for linking UPFs. The ANS_USE_UPF environment variable should not be set when using other methods to link UPFs.

To use this method start Mechanical APDL in batch mode and include one or more /UPF commands in the specified input listing. When the program reads the input and detects /UPF, the appropriate DLL is created.

You can include /UPF anywhere in your input file, and you can repeat /UPF as many times as needed to include multiple user routines in the relinked version. The following user routines can be linked using this method:

UANBEG

UANFIN

UCNVRG

UELMATX

UITBEG

UITFIN

ULDBEG

ULDFIN

USER01 -USER10

USERCNPROP

USERCREEP

USERCV

USERCZM

USERELEM

USERFLD

USERFRIC

USERFX

USERHYPER

USERINTER

USERMAT

USERMATTH

USEROU

USERSWSTRAIN

USER_TBELASTIC

USERWEAR

USOLBEG

USOLFIN

USRCAL

USREFL

USRSHIFT

USRSURF116

USSBEG

USSFIN

UTIMEINC

If you are running in a distributed memory parallel environment (Distributed ANSYS), you must set the ANS_USER_PATH environment variable to the working directory and include the name of the head compute node:

```
set ANS_USER_PATH=\\headnode\workingdirectory
```

When you run a user-linked version of the program by this method, the output includes this message:

```
NOTE - This ANSYS version was linked by /UPF with n user supplied routine(s).
```

where n indicates the number of routines specified by **/UPF** commands. The routines that have been linked will be included in the output listing.

Example Using Mixed Languages

The following directory contains an example of using the /UPF command method to link user routines that are written in mixed languages (FORTRAN, C, C++):

```
Program Files\ANSYS Inc\V211\ansys\custom\user\<platform>\Examples
```

The README.txt file in this directory contains complete instructions on running this example. This is a simple, automated test that verifies whether compilers are correctly installed and configured.

1.11.1.1. Using the /UPF Command on a Windows HPC Server System

Running a distributed memory parallel (Distributed ANSYS) solution on a Windows HPC Server system is more involved than running across a simple Windows cluster. You can use the steps de-

scribed here to test the /UPF command on a Windows HPC Server system. Several files are included with the ANSYS, Inc. software installation for use in this example.

Before you begin, Visual Studio 2017 Professional and Intel FORTRAN must be on the head compute node.

The files needed for this example can be found in the following directory:

```
C:\Program Files\ANSYS Inc\v211\commonfiles\MPI\WindowsHPC\UPF
```

These include two user routines, a Mechanical APDL input file, and two files required by the HPC Job Manager:

```
usercreep.F
usermat.F
slupf.inp
RUNANSYS_UPF.xml
runansysupf.bat
```

Copy these files to:

```
C:\Temp\%USERNAME%
```

You will need to modify the value of the **ANS_USER_PATH** environment variable in RUN-ANSYS_UPF.xml to the appropriate location where the UPF DLL library resides.

Submit RUNANSYS_UPF.xml to the HPC Job Manager. When the job is complete, run the following commands from a Command Prompt window:

```
clusrun /exclude:%CCP_SCHEDULER% copy /y C:\Temp\%USERNAME%\Work\*.out \\%CCP_SCHEDULER%\Temp\%USERNAME%
findstr /I debug *.out
```

If the test worked correctly, this should display many lines of debug from all output files.

For more information on running Distributed ANSYS under Windows HPC Server, see Configuring Distributed ANSYS in the *Parallel Processing Guide*.

1.11.2. Creating a Dynamic-link (DLL) Library

You can also set up UPFs on Windows systems by using a DLL library. All FORTRAN files (files ending with .F), C files (files ending with .c), and C++ files (files ending in .cpp) that you want to include in your DLL library should reside in your working directory. To compile all *.F, *.c, and *.cpp routines, issue the following command:

```
\Program Files\Ansys Inc\v211\ansys\custom\user\<platform>\ANSUSERSHARED.bat
```

After you issue ANSUSERSHARED. bat, the output will display the options for building the DLL library. The first portion of the output is shown below:

```
This is the ANSYS 2021 R1 ANSUSERSHARED script. It is used to build a DLL of User Programmable Features for the program.
```

```
NOTE: The user subroutine source file(s) are expected to reside in this directory.

Enter one of the following choices to create your User Programmable Feature DLL:
```

The user routines that are supported by this method will appear in a list. (These are the same user routines as listed above for the /UPF command method (p. 140).) Enter the user routine name you wish to include. As an example, if you enter USERMAT the following output will display and prompt you to select another option:

```
You chose USERMAT
Microsoft (R) Incremental Linker Version 14.10.25027.0
Copyright (C) Microsoft Corporation. All rights reserved.
-out:UserMatLib.dll
-def:UserMatLibex.def
-411
-machine: AMD64
-map
-manifest
-manifestfile:UserMatLib.dll.intermediate.manifest
-defaultlib: ANSYS.lib
-defaultlib:bufferoverflowU.lib
  Creating library UserMatLib.lib and object UserMatLib.exp
     UserMatLib.dll has been successfully built.
     Set the environment variable ANS_USER_PATH to the directory where the
     UserMatLib.dll resides and run ansys211 to use your newly generated
     user shared library.
     *****************
```

After you have selected all desired user routines, press **Enter** to quit.

If you are running in a distributed memory parallel environment (Distributed ANSYS), you need to include the name of the head compute node when specifying the working directory:

```
set ANS_USER_PATH=\\headnode\workingdirectory
```

When you run a user-linked version of Mechanical APDL, the output echos the value of ANS_USER_PATH and will include the following:

```
NOTE: This Mechanical APDL version was linked by Licensee
```

To return to the original version of Mechanical APDL, return the **ANS_USER_PATH** environment variable to its original value.

Multiple UPF DLLs can be created via the ANSUSERSHARED. bat script but must exist in the same directory as designated by the ANS_USER_PATH environment variable.

1.11.3. Using the ANS_ADMIN Utility

The **ANS_ADMIN** procedure for compiling and linking UPFs on Windows Systems creates a custom executable. This executable can be used to run in a shared memory parallel (SMP) environment or a distributed memory parallel environment (Distributed ANSYS).

As mentioned previously, the source files for the user routines reside in subdirectory Program Files\ANSYS Inc\V211\ansys\customize\user\.

If you modify any of the user routines, move them to the folder(s) they are linking in. By default this folder is:

Program Files\Ansys Inc\V211\ansys\custom\user\<platform>

Where <platform> is a directory that uniquely identifies the hardware platform version: "Winx64"
for 64-bit Windows.

Note:

The user routines listed in Using the /UPF Command (p. 140) cannot be linked with the ANS_ADMIN utility or the ANSCUST. bat script. These user routines have reserved DLL names and must be built using either the /UPF command or the ANSUSERSHARED. bat script.

You can select the **Relink ANSYS** option from the **ANS_ADMIN211** utility to link these changes into Mechanical APDL. This procedure will ask which versions you intend to relink and then will compile all FORTRAN files (files ending with .F), C files (files ending with .c), and C++ files (files ending in .cpp) in the Program Files\ANSYS Inc\V211\ansys\custom\user\platform> directory. The procedure then loads all object files (files ending with .obj), along with the default Mechanical APDL objects and libraries and creates custom executables. The executable file(s) created will be named ansys.exe and will reside in the folders described above.

Caution:

When creating custom executables, they must be named ansys.exe. This requirement is due to shared library usage.

Note:

On any Windows system, if you are attempting to create a relinked version of Mechanical APDL by using ANSCUST instead of using the **ANS_ADMIN211** utility (as recommended above), error messages may occur. These messages may state that object files have been created, but the Mechanical APDL executable has not been created; or the errors may state that some libraries necessary to complete the link cannot be found. These errors usually indicate that required compiler environment variables are not set. To avoid these errors, use the following workaround when relinking Mechanical APDL with ANSCUST:

 Pick Start > All apps > Visual Studio 2017 > Developer Command Prompt for VS2017, which should open a new command prompt window.

- In this command prompt window, issue the drive letter and all the necessary cd commands to move to the directory where the customized files exist (example: C:\Program Files\ANSYS Inc\V211\ansys\custom\user\).
- Type ANSCUST in this command window. The process of creating the new usercustomized Mechanical APDL version begins.

After relinking the Mechanical APDL executable, the program can be executed by either of the following two methods:

- 1. To execute the relinked version of the Mechanical APDL program:
 - Click Start>Programs>ANSYS 2021 R1>Mechanical APDL Product Launcher
 - In the launcher, select the **Customization/Preferences** tab, then browse to the path which contains the relinked ansys.exe. Select other desired options then pick **Run** to execute the customized ansys.exe.
- 2. To execute the relinked ansys.exe from a Command Prompt window, use one of the following commands.
 - · Interactive:

```
ansys211 -p roduct variable> -g -custom <path>
```

Batch:

```
ansys211 -b -p roduct variable> -j jobname -i <input file> -o <output file> -custom <path>
```

where "path" indicates the full path to the relinked ansys.exe.

Note:

The -custom option must be the last option at the end of the command line.

Note:

Output from a user-linked version contains the following statement:

This Mechanical APDL version was linked by Licensee

1.12. Activating UPFs

The Mechanical APDL program activates many UPFs via a specific user action. This can be through a command option or a user selection. Below is a list of specific actions required for several types of UPF.

 To activate user elements created using the method described in Creating a New Element via the User-Defined Element API (p. 151), you need **USRELEM** and **USRDOF** commands, as well as ET and TYPE commands.

- To activate a user element created using the method described in Creating a New Element by
 Directly Accessing the Program Database (p. 169), you must select it as one of the element types
 in a model using the ET command, then set the element attribute pointer using the TYPE
 command, and define elements using the solid modeling or direct generation method.
- To define a user material described in Subroutine UserMat (Creating Your Own Material Model) (p. 199), Subroutine UserCreep (Defining Creep Material Behavior) (p. 221), and Subroutine userswstrain (Defining Your Own Swelling Laws) (p. 228), you need to activate it with the corresponding TB commands.
- To customize contact interfacial behaviors as described in Subroutine USERFRIC (Writing Your Own Friction Laws) (p. 236) and Subroutine USERINTER (Writing Your Own Contact Interactions) (p. 238), you need to activate them with the corresponding **TB** commands.
- To program history-dependent contact properties described in Subroutine USERCNPROP (Defining Your Own Real Constants for Contact Elements) (p. 232), you need to activate the user routine with the **R**, **RMORE**, or **RMODIF** command. The real constant must be defined by the Mechanical APDL reserved table name CNPROP and enclosed in % signs (that is, % CNPROP%).

UPFs that are not activated by the means described above must be activated by either of the following methods:

- · Issuing the USRCAL command
- Choosing menu path Main Menu>Preprocessor>Loads>-Load Step Opts->Other>User Routines or Main Menu>Solution>-Load Step Opts->Other>User Routines.

To activate or deactivate the routines, issue the command **USRCAL**, Rnam1, ...Rnam9, where Rnam1 and Rnam9 are the names of specific routines. You can specify up to nine routines with one **USRCAL** command, or you can issue multiple **USRCAL** commands.

Issue the command **USRCAL**,NONE to deactivate all valid user subroutines. To list the status of the routines, issue the command **USRCAL**,STAT.

For a list of the user routines that the **USRCAL** command (or its equivalent menu paths) affects, see the **USRCAL** command description in the *Command Reference*.

If you do not activate the UPFs in this manner, standard Mechanical APDL logic is used by default. For example, when you apply a convection load, standard Mechanical APDL logic is the default even if you have a user convection routine linked in. The user convection routine must be activated by the **USRCAL** command or its menu equivalent.

1.13. Running Your Custom Executable

You can run a custom executable from the **Customization/Preferences** tab of the launcher:

Enter the full pathname to the custom executable in the **ANSYS Custom Executable** field. Do not include the -custom argument.

When run from the command prompt, if no path is specified after the -custom argument, the ansys211 script searches the current working directory for the custom Mechanical APDL executable (ansyscust.e by default on Linux or ansys.exe on Windows). If the custom Mechanical APDL ex-

ecutable resides in a separate directory (or has a name other than ansyscust.e on Linux), you can specify a different path and filename after the -custom argument.

Caution:

If you are running on a Windows system and you create a custom Mechanical APDL executable, it must be named ansys.exe. This requirement is due to shared library usage.

On Linux, you can also run your custom executable via command line.

ansys211 -custom /pathname/ansyscust.e

1.14. Verifying Your Routines

After compiling and linking your new user routine, test and verify it using whatever procedures you think are adequate. Remember, verifying that your customized version of Mechanical APDL works properly is *your* responsibility.

Make certain that your custom version of the program performs correctly for the combinations of elements, analysis types, materials, boundary conditions, and so on that you plan to use. Confirm that the logic you introduced is correct and does not produce any unwanted side effects.

In testing your custom user routines, you also should verify that the changes you have made do not affect standard, non-customized Mechanical APDL features. To do so, you can compare the results of a set of problems from the *Mechanical APDL Verification Manual* run on the standard version and on the customized version. Input for these problems is also available on your product distribution medium.

Always remember: your last step, a series of steps, or even your concept may be wrong. Proceed in clear steps, and verify your work as often as possible. Keep intermediate versions of your modified source code on backup media.

Note:

If you contact your site's ANSYS system-support person or any ANSYS, Inc. representative about the performance of a custom version of Mechanical APDL, always tell that person that you are using a user programmable feature. If you feel that an error exists in an unrelated feature of Mechanical APDL, demonstrate the suspected error in a non-customized, production version of the program before you report the error to an ANSYS, Inc. representative.

1.15. Debugging Commands

To debug errors in your user routines, you can use commands not documented in the *Command Reference*. Use these commands only for extremely small models with few solution iterations (otherwise, they will generate an excessive amount of output).

Two useful commands are **OUTEQ** and **/NERR**. The command **OUTEQ**,ON can be used to output results from all equilibrium iterations. The command **/NERR**,,,,-1 causes errors to be reported as before, but the run continues anyway, normally terminating with either a) system abort or b) incorrect answers. The

/NERR,,,-1 command is intended for program debugging and may generate erroneous results. You should remove this statement before generating solutions for production use.

1.16. Other Useful Commands

Two other Mechanical APDL commands, **NSVR** and **/UCMD**, can help you implement UPFs. (Neither command has an equivalent GUI path.) Use the **NSVR** command to define the number of extra variables that need to be saved for user programmable element options, such as user plasticity.

Issue the **/UCMD** command to make a user routine into a custom command. For more information, see Defining Your Own Commands (p. 261).

1.17. Generating Output

You can generate output controlled by the **/OUTPUT** command by using the FORTRAN write statement. The output unit for this statement is usually called IOTT. IOTT may be defined with the function wringr. See the discussion on the function wringr in Subroutines for Your Convenience (p. 343) for more details.

1.18. Reading Large Data Files More Rapidly

When files containing Mechanical APDL-related data are large, loading them into the program or writing them out to an external file can be a slow process. For example, consider a problem file which contains nearly 462,000 lines, 150,000 of which contain nodal data and 97,383 of them containing data for elements. Because many of the lines in this file are in command format, Mechanical APDL requires much time to read it.

You can shorten the time the program requires to read such files by including two commands in your programs, UPFs, or macros: EBLOCK (p. 113) and NBLOCK (p. 117). The NBLOCK (p. 117) command converts nodal data into fixed format data blocks (which Mechanical APDL can read more quickly than commands). The EBLOCK (p. 113) command places element data into a fixed format block, one line per element. These commands also compress displacement constraint data to one line per constraint node. See The CDWRITE (CDB) File Format (p. 99) in the *Guide to Interfacing with ANSYS* for more information on the use of these commands.

Chapter 2: UPF Subroutines and Functions

This guide describes the various subroutines, functions, and commands that allow you to customize the program for your specific purpose. The first portion of each subroutine or function (consisting of comment lines) is shown in most cases.

User subroutines that can be modified have the term *USERDISTRIB* in the first line of the subroutine. For example, the first line of the usercnprop subroutine looks like this:

*deck,usercnprop

USERDISTRIB

User subroutines that do not have *USERDISTRIB* in the first line cannot be modified and must be used as they are.

The following UPF topics are available:

- 2.1. Creating a New Element
- 2.2. Supporting Subroutines for Element Creation
- 2.3. Subroutines for Modifying and Monitoring Existing Elements
- 2.4. Subroutines for Customizing Material Behavior
- 2.5. Subroutines for Customizing Contact Interfacial Behavior
- 2.6. Subroutines for Customizing Loads
- 2.7. Subroutines for Sharing Data Between User Routines
- 2.8. Running Mechanical APDL as a Subroutine
- 2.9. Defining Your Own Commands
- 2.10. Support Subroutines
- 2.11. Access at the Beginning and End of Various Operations
- 2.12. Memory-Management Subroutines
- 2.13. Parameter-Processing Subroutines
- 2.14. Other Useful Functions

2.1. Creating a New Element

Two tools are available for creating a user-defined element:

- The user-defined element API (p. 151)
- Direct access to the program database and files (p. 169)

ANSYS, Inc. recommends the user-defined element API in most cases. The direct-access method is generally for special-purpose use only, or if you are already using preexisting elements created with this method.

This table highlights the differences between the two methods:

Interface	User-defined element API	Accessing program database and files directly
Description	Offers a simpler interface while preserving much of the underlying user-element capability. An understanding of the database subroutines and the file structure is rarely necessary to use the interface.	No special interface. With few exceptions, if a capability exists for an element, it will exist here. The logic necessary for using this interface effectively is more complex.
Relative level of difficulty	Medium	High
Expected compatibility between versions	High	Medium
Element names	USER300	USER100 to USER105
Demonstration logic included on your product distribution media	4-node quad and 20-node brick elements	uel100 (a structural mass element) and uel101 (a simple link element)
Typical linear material access subroutine	getMatProp	propev
New nonlinear material properties	Program in UserMatTh.	No special programming has been set up.
Existing nonlinear material properties	All standard structural materials are accessible via ElemGetMat.	Limited capability. Accessible via plastx, creepx, and swellx.
Non-structural material properties	No special programming has been implemented.	No special programming has been implemented.
Number of different element types allowed	Effectively, no limit.	Effectively, no limit.
Element characteristic capability	Input the basic 10 element characteristics (via the USRELEM and USRDOF commands). All other characteristics default automatically.	Input all 140 element characteristics (using uec100). Approximately 30 are normally used, and the rest default unless required for special cases.
Applicable subroutines to be programmed	UserElem	uec100, uel100, and rarely uex100 and uep100. Subroutines uec101 to uec105 are analogous.
Access to database information	Generally through the interface.	Through subroutines (such as tmp-get, prsget, svgidx, svrget, svpidx, svrput).
Access to file information	Through the interface.	Through pointers and subroutines (such as eldwrtL, eldwrnL).
Special features	Element convergence criteria Cutback control via element	None.
Capabilities <i>not</i> included	Control information unavailable for:	Material TB labels PLASTIC, NLISO, AHYPER, HYPER, PRONY, SHIFT,

Interface	User-defined element API	Accessing program database and files directly
	Birth and death Superelement stress pass Initial stress Section input Input of fluences Swelling	ELASTIC, ANEL, SDAMP, SMA, CAST, EDP, and GURSON.

2.1.1. Input and Output Abbreviations

The descriptions of the subroutines or functions within this chapter describe both the input arguments and output arguments. Argument information includes the argument's type, size and intent.

• Argument *type* is one of the following:

int - integer

dp - double-precision

log - logical

chr - character

dcp - double-precision complex

• Argument size is one of the following:

sc - scalar variable

ar(n) - array variable of length n

func - functional return value

• Argument *intent* is one of the following:

in - input argument

out - output argument

inout - both an input and an output argument

2.1.2. Creating a New Element via the User-Defined Element API

Following is the general process for creating your own element via the user-defined element API.

Steps 2 and 3 specify data for the user-defined element API. All other steps represent standard features.

Step Description		Comments	
1.	Specify the element	Issue the ET and TYPE commands. The name of the element must be	
	type.	USER300.	

2.	Define your new element according to the specified element type.	Issue the USRELEM command. Specify the element characteristics (such as the number of nodes, number of dimensions, number of real constants etc.).
3.	Specify nodal degrees of freedom.	Issue the USRDOF command. You can specify a maximum of 10 degrees of freedom per USRDOF command; to define additional degrees of freedom, issue the command again. Each node will have the same degrees of freedom. Although you can
		specify any valid degrees of freedom, the total number of degrees of freedom for your element cannot exceed 480, and the number of degrees of freedom for each node cannot exceed 32.
4.	Define real constants.	If needed.
5.	Create finite element models.	 Use either of these methods: Direct generation Create elements directly from nodes, using commands such as E, EGEN, EN, ENGEN, or EMORE. (You can also use the CDREAD command if the .cdb file is available.) This method is the only way to create an element with a topology different from that of any standard element. Meshing commands This method is available only if your element has the same topology as that of a standard element and you have specified any standard element shape (USRELEM KeyShape value) except ANYSHAPE.
6.	Apply boundary conditions and loads.	As needed.
7.	Specify solution options.	If your element has multi-field degrees of freedom (displacements and temperatures).
8.	Perform postprocessing.	Postprocessing occurs normally as with any other element. Only total strain (or equivalent quantities such as thermal gradient) and stress (or equivalent quantities such as thermal flux) are saved as regular result quantities. Other variables are saved as nonsummable miscellaneous variables in the results file.

Recommendations and Restrictions

The following recommendations and restrictions apply to user-defined element USER300:

- Verify that your input data for the **USRELEM** and **USRDOF** commands are consistent with the values used in the <code>UserElem.F</code> code. For example, if the number of dimensions (*NDIM*) specified via the **USRELEM** command is 2, do not change the number of dimensions specified in the <code>UserE-lem.F</code> subroutine from 2. A runtime error or incorrect results can occur if the values do not match.
- The program may activate default solution settings automatically according to the USER300 element's degrees of freedom, but the default solution control settings may not be optimal for your element.

• The USER300 element does not support section (SECxxx) commands. For composite beams and layered shells, you must input element data via real constants and code the UserElem. F subroutine accordingly.

2.1.2.1. Subroutine UserElem (Writing Your Own Elements)

The UserElem subroutine provides an interface to program code above the element level. UserElem supports shared memory and distributed parallel processing; however, you are responsible for ensuring that your code can use parallel processing.

The subroutine passes all data needed to create a user-defined element and returns all data and results from the element to update the program database and files. With this API, you can create virtually any element type *without* having to access program database and files directly (p. 169). Two examples are included in this subroutine: a 4-node quadrilateral 2-D element, and 20-node brick structural element, both for geometric linear analysis. Key options (KEYOPT settings) switch the elements.

The following table shows the input and output arguments, and their definition and usage. Some argument names (such as those pertaining to element matrices and load vectors) are common to structural analyses; however, you can specify argument values appropriate to analyses in other engineering disciplines.

Argument	Input (I) or Output (O)	Definition	Purpose	How Defined
elId	I	Element number	Output information	At FE model creation
matId	I	Material number	Output information Call material subroutines	At FE model creation
keyMtx	I	Formulation request	Specifying which matrices and load vectors to form	Program code
lumpm	I	Mass matrix format: 0 = Consistent mass matrix 1 = Lumped mass matrix	Specifying how to form the mass matrix	LUMPM command
nDim	I	Number of dimensions	Element coding	USRELEM command
nNodes	I	Number of element nodes	Element coding	USRELEM command
Nodes	I	Element node list Connectivity	Output	At FE model creation

nIntPnts	I	Maximum number of element integration points	Element coding	USRELEM command
nUsrDof	I	Number of element degrees of freedom	Element coding The degrees of freedom are ordered in the way in which they are listed via the USRDOF command for each node and repeated for all nodes All element matrices DOF values and load vectors must be arranged in the same way	USRELEM and USRDOF commands
kEStress	I	Element stress state	Element coding Calling material subroutines if requested	USRELEM command
keyAnsMat	I	Element formulation key: 0 Write your own material formulation 1 Use standard material subroutines and call ElemGetMat subroutine	Specifying how to create material data	USRELEM command

keySym	1	Flag for symmetricity of element matrices	Element coding Program assembly logic	USRELEM command
nKeyOpt	I	Maximum number of element key options allowed (up to 2)	Element coding	Program code
KeyOpt	I	Element key options KEYOPT(1) = 0~99	Branching the user-element codes to different formulations. (This could be equivalent to 100 x 100 different types of elements.)	ET command
temper	I	Nodal temperatures at current time	Temperature dependence and thermal loads	BF and BFE commands (if key-Shape is specified in the UserE-lem subroutine)
temperB	1	Nodal temperatures at the end of the last substep	Temperature dependence and thermal loads	Program code
tRef	I	Reference temperature	Temperature dependence and thermal loads	TREF command
kTherm	I	Key indicating whether a thermal load exists: 1 = Calculate the thermal load 0 = No thermal load calculation	Element coding	
nPress	I	Number of pressure values	Element coding The size of the press vector	USRELEM command At FE model creation

		Ť		ĺ
Press		Pressures at nodes of element facets (available only when keyShape is specified via the USRELEM command) The pressure vector is ordered in the element with the same topology as that in the standard element library. Refer to that element for details.	Pressure load and pressure load stiffness	SF and SFE commands
kPress	I	Key indicating whether a pressure load exists: 1 = Calculate the pressure load	Element coding	Program code
		0 = No pressure load calculation		
nReal	I	Number of real constants	Element coding	USRELEM command
RealConst		The list of real constants	Element coding Can pass material properties, section/layer information, element material control, and any element data	R command
nSaveVars	I	The number of variables saved in the .esav file for the element	Element coding The size of saveVars	USRELEM command
saveVars	I/O	The data saved in the .esav file The program saves the data after	Element coding	UserElem subroutine

	Υ			1
		exiting the UserElem subroutine and retrieves it immediately before entering UserElem again. It should include kinematic related variables only when the material subroutine is called; otherwise, it should include both kinematic and material data. History dependent variables can only be saved/updated when the substep is converged (key- HisUpd = 1).		
xRef	I	Initial coordinates of the element nodes Values in global Cartesian coordinates	Element coding	At FE model creation
xCur	I	Current (deformed) coordinates of element nodes Values in global Cartesian coordinate system, equal to xRef when nlgeom = off	Element coding	Program code
TotVal- Dofs	I	Total values of degrees of freedom (displacements for structural analysis) Values in global Cartesian coordinate system	Element coding	Program code
IncVal- Dofs	I	Increment values of degrees of	Element coding	Program code

		1		
		freedom occurring at the current substeps Values in global Cartesian coordinate system		
ItrVal- Dofs	I	Iteration values of degrees of freedom occurring at the last iteration	Element coding	Program code
		Values in global Cartesian coordinate system		
VelVal- Dofs	I	First time derivatives of degrees of freedom	Velocities	Program code
AccVal- Dofs	I	Second time derivatives of degrees of freedom	Accelerations	Program code
kfstps	I	Key indicating first time entering the element subroutine:	Initializing data	Program code
		1 = First time 0 = Other than first time		
nlgeom	I	Flag indicating whether large displacement/deformation is in effect	Element coding	NLGEOM command
nrkey	I	Newton-Raphson algorithm key: 1 Any nonlinear analysis 0 Pure linear analysis	Output Element coding	
outkey	I	Key indicating output result type: 1 This is an output call, the substep is converged, and you can print/save element results 0 All other cases	Element coding	Program code

iott	I	Key indicating whether any element output should appear in the print file: 0 = No 1 = Yes Output file number	The FORTRAN output file number. All information written in the specified file appears in the output file.	Program code OUTPR command Program code
keyHisUpd	I	Key to update history-dependent variables: 1 = The substep converged; ready to update history-dependent variables (such as equivalent plastic strain) 0 = Solution not yet converged; cannot update history-dependent variables	Element coding	Program
The following You can usual		for debug, timing, and conve	ergence-control purpo	oses only.
ldstep		Current load step number	Output Debug	Program code
isubst	I	Current substep number	Output	Program code
ieqitr	I	Current iteration number	Output	Program code
timval	I	Current time	Output	Program code
keyEleErr	I/O	Formulation error key: 0 = No error (preset value) 1 = Error occurred in element	Element coding	Program code

		formulation, possibly due to excessive deformation. (The program lessens deformation if possible by cutback.)				
keyEleCnv	I/O	Element convergence key: 1 = Converged (preset value before calling) 0 = Not converged	Provides manual control of convergence when you introduce any constraint at the element level (such as volumetric constraint for mixed u-P)	Program code		
End of special	End of special-purpose variable group					
eStiff	0	Small-deformation stiffness matrix In global Cartesian coordinate system	Solution	Requested when keyMtx(1) = 1		
eMass	0	Mass matrix In global Cartesian coordinate system	Solution	Requested when keyMtx(2) = 1		
eDamp	0	Damping matrix In global Cartesian coordinate system	Solution	Requested when keyMtx(3) = 1		
esstiff	0	Stress stiffness matrix In global Cartesian coordinate system	Solution	Requested when keyMtx(4) = 1		
fExt	0	External load vector In global Cartesian coordinate system	Solution	Requested when keyMtx(5) = 1		
fInt	0	Internal nodal force vector In global Cartesian coordinate system	Solution	Requested when keyMtx(6) = 1		
elVol	0	Element volume	Output	UserElem subroutine		

elMass	0	Element mass	Output	UserElem subroutine
elCG	0	Element centroid coordinates In global Cartesian coordinate system	Postprocessing	UserElem subroutine
nRsltBsc	I	Number of basic result data saved in result file	Specifying the size of RsltBsc	Program code
RsltBsc	0	Basic result data saved in results file These variables are accessible via the PRESOL and PRNSOL commands in the standard way and can also be plotted if you specify a KeyShape value via the USRELEM command.	Postprocessing	UserElem subroutine
nRsltVar	I	The number of result data to be saved in the result file as non-summable miscellaneous variables	Specifying the size of RsltVar	USRELEM command
RsltVar	0	The result data saved in the result file as non-summable miscellaneous variables The data is accessible via the PLESOL command only, but only one value for an element each time	Postprocessing	UserElem subroutine
nElEng	I	Number of energies Fixed at 3	Solution	UserElem subroutine
elEnergy	O	Element energy vector: elEnergy(1) Strain energy	Output	UserElem subroutine

	elEnergy(2) Plastic energy	
	<pre>elEnergy(3) Creep energy</pre>	

```
*deck,UserElem
                                    USERDISTRIB
c Copyright ANSYS. All Rights Reserved.
      subroutine UserElem (elId, matId, keyMtx, lumpm, nDim, nNodes,
     &
                           Nodes, nIntPnts, nUsrDof, kEStress,
     &
                           keyAnsMat, keySym, nKeyOpt, KeyOpt,
                           temper, temperB, tRef, kTherm,
     &
                           nPress, Press, kPress, nReal, RealConst,
     &
                           nSaveVars, saveVars, xRef, xCur,
                           TotValDofs, IncValDofs, ItrValDofs,
                           VelValDofs, AccValDofs,
                           kfstps, nlgeom, nrkey, outkey, elPrint, iott,
                           keyHisUpd, ldstep, isubst, ieqitr, timval,
                           keyEleErr, keyEleCnv,
                           eStiff, eMass, eDamp, eSStiff,
                           fExt, fInt, elVol, elMass, elCG,
                           nRsltBsc, RsltBsc, nRsltVar, RsltVar,
                           nElEng, elEnergy)
                 ***************
c *** Primary function: General User Element Subroutine
c *** Note:
        This routine is completed with an example, see more details later.
C
С
С
      PROGRAMMER SHOULD NOT CHANGE ANY PURE INPUT ARGUMENTS (marked by ....,in)!
С
С
      elId
                (int,sc,in)
                                   element number
С
С
      matId
                (int,sc,in)
                                   material number of this element
С
     keyMtx
                (int,ar(10),in)
                                   matrix and load vector form requests
                                      0 = not requested, 1 = requested
C
С
                                      see below for more details
С
      lumpm
                (int,sc,in)
                                   mass matrix format
                                     = 0 no lumped mass matrix
С
                                     = 1 lumped mass matrix
С
С
     nDim
                (int,sc,in)
                                   number of dimensions of the problem
                                        (defined on USRELEM command as NDIM)
С
С
                                     = 2.2D
                                     = 3 3D
C
С
      nNodes
                (int,sc,in)
                                   number of nodes of the element
                                        (defined on USRELEM command as NNODES)
C
                (int,ar(nNodes),in)node list of this element
      Nodes
C
С
      nIntPnts
                (int,sc,in)
                                   maximum number of integration points
С
                                        (defined on USRELEM command as NINTPNTS)
     nUsrDof
                                   number of DOFs of this element (matrix and
С
                (int.sc.in)
                                      load vector size)
С
С
     kEStress (int,sc,in)
                                   kEStress
                                        (defined on USRELEM command as KESTRESS)
C
С
      keyAnsMat (int,sc,in)
                                   key to indicate if ANSYS material
                                      routine is going to be called
C
С
                                      (defined on USRELEM command as KEYANSMAT)
С
                                      = 0, No
                                      = 1, Yes
C
                                   key to indicate if element matrices
С
      keySym
                (int,sc,in)
С
                                      is symmetric
С
                                        (defined on USRELEM command as KEYSYM)
С
                                      = 0, symmetric
С
                                      = 1, unsymmetric
С
      nKeyOpt
                (int,sc,in)
                                   number of element key options able to be
С
                                      used in this routine
С
      KeyOpt
                (int,ar(nKeyOpt),in) values of element key option defined
                                      by et or keyopt command for the
```

```
С
                                       user elements, only the first
                                       nKeyOpt values are passed in and can
С
                                       be used to branch the routine for
С
С
                                       different formulations
С
      temper
                 (dp,ar(nNodes),in) nodal temperatures at current time
                 (dp,ar(nNodes),in) nodal temperatures at the beginning of this
С
      temperB
                                       incremental step (substep)
С
                                    reference temperature
C
      tRef
                 (dp,sc,in)
      kTherm
                 (int,sc,inout)
                                    input: flag for thermal loading
C
C
                                        = 1, Temperatures at nodes are different
                                        from the reference temperature,
C
                                        thermal loading might be needed.
С
С
                                        = 0, Temperatures at nodes are the same
                                        as the reference temperature,
C
                                        thermal loading is not needed.
С
                                    output: flag for thermal strains
С
                                    number of pressure values for this element
С
      nPress
                 (int,sc,in)
                 (dp,ar(nPress),in) applied elemental face load (pressure)
С
      Press
C
      kPress
                 (int,sc,in)
                                    flag for pressure loading
С
                                        = 1, pressure load is applied and
                                        equivalent nodal forces should be
C
                                        calculated
C
С
                                        = 0, no pressure loading
С
      nReal
                 (int,sc,in)
                                    number of real constants
                                         (defined on USRELEM command as NREAL)
C
                                    user defined real constants
С
      RealConst (dp,ar(nReal),in)
                                    number of saved variables
С
      nSaveVars (int,sc,in)
C
                                        (defined on USRELEM command as NSAVEVARS)
                (dp,ar(nSaveVars),inout) user saved variables
С
      saveVars
                 (dp,ar(nDim,nNodes),in)
С
      xRef
С
                                    nodal coordinates in initial configuration
С
      xCur
                 (dp,ar(nDim,nNodes),in)
С
                                    nodal coordinates in current configuration
С
      TotValDofs (dp,ar(nUsrDof),in) total values of DOFs (displacements)
С
                                       from time = 0
      IncValDofs (dp,ar(nUsrDof),in) incremental values of DOFs (displacements)
C
С
                                       for the current step
С
      ItrValDofs (dp,ar(nUsrDof),in) iterative values of DOFs (displacements)
                                       for the current iteration
С
С
                                       (normally needed for debug only)
      VelValDofs (dp,ar(nUsrDof),in) first time derivatives of DOFs
С
С
                                                (velocities) (normally not needed)
      AccValDofs (dp,ar(nUsrDof),in) second time derivatives of DOFs
С
С
                                             (accelerations) (normally not needed)
                                    key for the first iteration of first
С
      kfstps
                 (int,sc,in)
С
                                       substep of the first load step
                                       = 1 ves
C
С
                                       = 0 no
С
      nlgeom
                 (int,sc,in)
                                    large deformation key [from nlgeom command]
                                       = 0 NLGEOM, OFF
С
                                       = 1 NLGEOM, ON
С
С
      nrkey
                 (int,sc,in)
                                    key to indicate a newton-raphson
С
                                       (incremental) procedure
                                       = 0 No
С
                                       = 1 Yes
C
С
      outkey
                 (int,sc,in)
                                    key to indicate if any element output is
С
                                       to be placed on the print file or the
                                       result file
C
С
                                       = 0 No
С
                                       = 1 Yes
                                    key to indicate if any element output is
      elPrint
                (int,sc,in)
C
С
                                       to be placed on the print file
                                       = 0 No
С
                                       = 1 Yes
C
С
      iott
                 (int,sc,in)
                                    print output file unit number
                                    key to indicate if history-dependent
      keyHisUpd (int,sc,in)
C
С
                                      variables need to be updated, like
С
                                      equivalent plastic strain, back stress
                                      etc. since the iteration is already
C
С
                                      converged
С
                                       = 0 not converged, don't need to update
```

```
С
                                           history dependent variables
С
                                       = 1 yes, converged, need to update
                                           history dependent variables
С
С
С
      --- The following 7 variable group can usually be ignored.
      --- The variables are used for debug, timing, and convergence control.
C
                                    current load step number
С
                (int,sc,in)
                                    current substep number
C
      isubst
                (int,sc,in)
                                    current equilibium iteration number
      iegitr
C
                (int.sc.in)
C
      timval
                (int,sc,in)
                                    current time value
                                    key to indicate if there is any element
      keyEleErr (int,sc,inout)
C
С
                                       formulation error, like negative Jacobian.
С
                                       The error could be caused by too
                                       large incremental step, illegal model.
C
С
                                       = 0 no error (preset value before calling)
С
                                       = 1 some error happens. ANSYS will
                                       decide to stop the analysis or cutback
С
                                       the substep (bi-section) based on other
С
C
                                       user input and information at higher
                                       level.
С
      keyEleCnv (int,sc,inout)
                                    key to flag if this element satisfies
C
                                       the user defined element convergence
C
С
                                       criterion.
                                       = 1, yes, the criterion is satisfied
С
                                         or don't have any criterion at all
C
С
                                         it is preset value before calling
С
                                       = 0, no, the element doesn't satisfy
                                         element convergence criterion. If
C
                                         this is the case, the iteration will
С
                                         not converge even when both force
C
                                         and displacement converge
С
        ---- end of 7 variable group -----
С
C
                                                                     requested if
C
      eStiff(dp,ar(nUsrDof,nUsrDof),inout) stiffness matrix
С
                                                                      keyMtx(1)=1
      eMass (dp,ar(nUsrDof,nUsrDof),inout) mass matrix
                                                                      keyMtx(2)=1
C
С
      eDamp (dp,ar(nUsrDof,nUsrDof),inout) damping matrix
                                                                      keyMtx(3)=1
С
      eSStiff(dp,ar(nUsrDof,nUsrDof),inout)stress stiffness matrix keyMtx(4)=1
      fExt.
                 (dp,ar(nUsrDof),out)
                                            applied f vector
                                                                      keyMtx(5)=1
C
С
      fInt
                 (dp,ar(nUsrDof),out)
                                            internal force vector
                                                                      keyMtx(6)=1
      elVol
                (dp,sc,out)
                                    element volume
С
      elMass
                                    element mass
С
                 (dp,sc,out)
С
      elCG
                 (dp,ar(3),out)
                                    element centroid coordinates in current
С
                                       configuration
С
      nRsltBsc
               (dp,sc,in)
                                    number of basic elemental results saved in
                                     result files
C
С
      RsltBsc
                (dp,ar(nRsltBsc),out) basic elemental results
С
                                         (see EXPLANATION below)
                                    number of elemental results saved in
      nRsltVar
                (int,sc,in)
C
                                       result file as non-summable miscellaneous
С
С
                                       variables
С
                                         (defined on USRELEM command as NRSLTVAR)
      RsltVar
                (dp,ar(nRsltVar),out) variables to saved in result files as
С
                                        non-summable miscellaneous variables
C
С
                                        requested when outkey = 1
С
C
      nElEng
                (int,sc,in)
                                    number of energies (fixed at 3)
С
С
      elEnergy (dp,ar(nElEng),out) elemental energy
                                       elEnergy(1), element strain energy
C
С
                                       elEnergy(2), element plastic strain energy
                                       elEnergy(3), element creep strain energy
С
C
      EXPLANATION OF RsltBsc
C
C
С
        Basic element results are saved and total number of result
С
      quantities is nRsltBsc, where:
             nRsltBsc = (7+7)* number of corner nodes at one element.
C
        To process the quantites by post processing properly, the results
C
      must be in the following order:
```

2.1.2.2. Subroutine ElemGetMat (Calling the Standard Structural Material Library)

The ElemGetMat subroutine is the API to the standard materials. When you issue the **USRELEM** command after setting the command's *KEYANSMAT* argument, the subroutine accesses the standard structural material library. It allows you to focus on the kinematic portion of element formulation while the program handles the material part of the formulation.

When calling the subroutine, input the associated material data via the standard method. There is no need to access this subroutine, only to call it.

The following table shows the input and output arguments, and their definition and usage.

Argument	Input (I) or Output (O)	Definition	Purpose	How Defined
elId	I	Element number	Output	At FE model creation
matId	I	Material number	Output information Getting material data	At FE model creation
nDim	I	Number of dimensions of element geometry 2 = 2-D element geometry 3 = 3-D element geometry Specifying the size of the nodal coordinates	Material calculation	At FE model creation
nTens	I	Number of stress/strain tensor components: 4 = 2-D and ordered as x, y, z, xy 6 = 3-D and ordered as x, y, z, xy, yz, xz	Specifying the data size	UserElem subroutine

nDirect	I	Number of direct component of stress/strain tensors	Specifying the data size	UserElem subroutine
		nDirect< or = nTens		
intPnt	I	Current integration point number	Output Data handling	UserElem subroutine
xCurIP	I	Coordinates of current integration point Values in global Cartesian coordinate system	Material calculation	UserElem subroutine
TemperIP	I	Integration point temperatures at the current time	Evaluating temperature-dependen material data	UserElem tsubroutine
TemperIPB	I	Integration point temperatures at the end of the last incremental step	Evaluating temperature-dependen material data	UserElem tsubroutine
IncStrain	I	Strain components [1] Incremental strain of the current substep when nlgeom = on Total strain when nlgeom = off	Material calculation	UserElem subroutine
defG0	I	Deformation gradient tensor at the end of previous substep [1]	Material updating	UserElem subroutine
defG	I/O	Total deformation gradient tensor at the current time [1]	The component in thickness direction is updated by material subroutines for plane stress and shell elements	UserElem subroutine
kTherm	I/O	Thermal loading key: 0 = No thermal loading 1 = Has thermal loading	Thermal load calculation	UserElem subroutine
cMat	0	Material Jacobian [1]	Forming stiffness	Material subroutine

MatProp	0	Material data for element formulation	Forming mass matrix Handling	Material subroutine
			transverse shear	
			Output	
Stress	0	Cauchy stress [1]	Forming geometric stiffness	Material subroutine
			Calculating internal forces	
Strain	0	Total strain components [1]	Output	Material subroutine
StressTh	0	Total thermal stress components [1]	Output Calculating thermal loading	Material subroutine
StrainTh	0	Total thermal strain components [1]	Output	Material subroutine
StrainPl	О	Total plastic strain components [1]	Output	
StrainCr	O	Total creep strain components [1]	Output	
StressBk	0	Backstress components [1]	Output	
StrainSw	0	Swelling strain	Not yet supported	
EnergyD	0	Energy density: 1 = Elastic energy density 2 = Plastic energy density 3 = Creep energy density		
MatRotGlb	O	Rotation matrix	Rotation matrix from global Cartesian to rotated element coordinate system	Used only for solid elements when nlgeom = on

^{1.} All tensor component values in the subroutine are in the global Cartesian coordinate system for solid elements, and in the co-rotated element Cartesian coordinate system for link, beam and shell elements.

```
*deck,ElemGetMat
     subroutine ElemGetMat (elId, matId, nDim, nTens, nDirect,
                                intPnt, xCurIP, TemperIP,
                                TemperIPB, kTherm, IncStrain,
    &
    &
                                defG0, defG, CMat, MatProp,
                                Stress, Strain, StressTh, StrainTh,
    &
                                StrainPl, StrainCr, StressBk, StrainSw,
    δ
                                EnergyD, MatRotGlb)
c *** Primary function: call ANSYS material routines to obtain material
                        data for USER300 elements
c *** Notice - This file contains ANSYS Confidential information ***
C
      input arguments
C
C
     -----
     elId
                                    element number
C
                 (int.sc)
     matId
                                    material number of this element
                 (int,sc)
C
С
     nDim
                 (int,sc)
                                    number of dimensions of the problem
                                    = 2 2D
С
                                    = 3 3D
C
                                    number of stress/strain components
С
     nTens
                 (int,sc)
     nDirect
С
                 (int,sc)
                                    number of stress/strain direct
C
                                      components
С
      intPnt
                 (int,sc)
                                    current integration point number
                                    coordinates of integration point
С
     xCurTP
                 (dp,ar(3))
С
     TemperIP
                 (dp,sc)
                                    integration point temperatures at
                                      current time
С
C
     TemperIPB
                 (dp,sc)
                                    integration point temperatures at
                                      the end of last incremetal step
С
С
      IncStrain
                 (dp,ar(nTens))
                                    strain for the current substep step when
                                       nlgeom = on
C
С
                                    total strain when nlgeom = off
С
     defG0
                 (dp,ar(3x3))
                                    deformation gradient tensor at the end
                                       of last incremental step
С
С
С
      input output arguments
                                    input desc
                                                  / output desc
      ========
                                                     ========
С
      defG
                                    deformation gradient tensor at current
С
                 (dp, ar(3x3))
С
                                      time, updated for thichness change in
С
                                      plane stress when nlgeom=on
С
     kTherm
                 (int,sc)
                                    flag for thermal loading
                                      input as:
C
С
                                      = 0 if temp = tref
С
                                      = 1 if temp .ne. tref
                                      gets reset to 0
C
                                                   if ALPX, ALPY, and ALPZ = 0
С
С
С
     output arguments
C
                 (nTens*nTens)
                                    material Jacobian matrix
C
     cMat
С
     MatProp
                 (dp,ar(5))
                                    commonly used materail properties
С
                                    MatProp(1), Gxz: shear modulus
                                    MatProp(2),Gyz: shear modulus
C
                                    MatProp(3), Gxy: shear modulus
С
С
                                    MatProp(4), density
                                    MatProp(5), nuxy
C
C
      Stress
                 (dp,ar(nTens))
                                    total stress
                                    total strain
С
     Strain
                 (dp,ar(nTens))
     StressTh
                 (dp,ar(nTens))
                                    thermal stress
C
     StrainTh
                 (dp,ar(nTens))
                                    thermal strain
C
     StrainPl
                 (dp,ar(nTens))
                                    plastic strain
С
С
     StrainCr
                 (dp,ar(nTens))
                                    creep strain
С
     StressBk
                  (dp,ar(nTens))
                                    back stress for kinematic hardening
     StrainSw
                                    isotropic swelling strain
С
                  (dp,sc)
С
                                        (swelling capability not available yet)
     EnergyD
                  (dp,ar(3))
                                    energy density
С
```

2.1.3. Creating a New Element by Directly Accessing the Program Database

The next few pages describe the user subroutines and supporting subroutines you use to create new elements. Using these subroutines, you can create new element types, add them to the element library, and use them as "regular" elements. You can create up to six independent element types (names USER100 - USER105). For demonstration purposes, the subroutines uel100 (for a structural mass element) and uel101 (for a simple link element) are included on the product distribution media.

2.1.3.1. User Subroutines

Subroutines uec100 through uec105 describe the element characteristics. Subroutine elccmt (on the distribution medium) describes the input for these subroutines in detail. You can use subroutines uex100 through uex105 to override default logic. Subroutines uec100 through uec105 define parameters such as:

- · 2-D or 3-D geometry
- Degree of freedom set
- Symmetric or unsymmetric matrix
- · Number of nodes
- Number of body loads (for example, temperatures)
- Number of surface loads (for example, pressures)
- · Number of real constants
- · Number of variables to be saved
- Number of rows in element matrices
- · Linear or nonlinear element.

Subroutines uel100 through uel105 calculate the element matrices (stiffness, specific heat, and so on), the element load vector (force, heat flow, and so on), and any element output quantities. The element printout also is generated, and the variables to be saved are calculated and stored in the results file.

Other user subroutines available for manipulating element information include the following:

• Subroutines uep100 through uep105 provide printed output of line elements. The general postprocessor, POST1, calls the subroutines, or you can call them using uel100 through uel105.

- Subroutine usertr allows access to the nodal transformations.
- Subroutine userac describes some of the data handling.

2.1.3.2. Subroutine uec100 (Defining Characteristics of the usr100 Subroutine)

```
*deck,uec100
     subroutine uec100 (elcdn,ielc,kerr)
      **** this subroutine defines the characteristics of user100.
C
С
         *** Copyright ANSYS. All Rights Reserved.
С
         *** ansys, inc.
C
c *** Notice - This file contains ANSYS Confidential information ***
С
     typ=int,dp,log,chr siz=sc,ar(n) intent=in,out,inout
C
С
c input arguments:
     variable (typ,siz,intent)
                                  description
C
     ielc (int,ar(IELCSZ),inout) - element characteristics
                                       see include deck elccmt for details
C
     kerr (int,sc,inout)
С
                                 - error flag up to this point.
C
                                    (do not initialize to zero)
C
c output arguments:
С
     variable (typ,siz,intent) description
     elcdn (chr,sc,out)
                                 - name of element
C
     ielc (int,ar(IELCSZ),inout) - element characteristics
С
С
                                         see include deck elccmt for details
     kerr (int,sc,inout)
                                - error flag (set to 1 if error)
С
С
      note to programmers: the validity of keyopt values may be checked here
С
```

2.1.3.2.1. Subroutines uec101 through uec105

The input and output arguments for subroutines uec101, uec102, uec103, uec104, and uec105 is identical to the uec100 subroutine listed above.

2.1.3.3. Subroutine uex100 (Overriding Element Characteristic Defaults)

```
*deck,uex100
                                  USERDISTRIB
     subroutine uex100 (ielc,kerr)
      *** subroutine to override element characteristic defaults ***
      *** hence, this routine is needed only in rare cases.
C
С
          *** Copyright ANSYS. All Rights Reserved.
          *** ansys, inc.
С
c *** Notice - This file contains ANSYS Confidential information ***
C
      *** input and output are the same as for uec100, except that this
С
      *** logic is called after the defaulting logic is finished.
С
      *** this defaulting is done in ansys subroutine echdft(not a upf).
С
      *** as indicated above, this routine is rarely needed, but if it is
C
      *** desired to see the effect of echdft, you may print out the ielc array
С
      *** leaving uec100 and print it out again entering this routine.
С
С
С
     typ=int,dp,log,chr siz=sc,ar(n) intent=in,out,inout
C
c input arguments:
     variable (typ,siz,intent)
С
                                    description
С
      ielc (int,ar(IELCSZ),inout) - element characteristics
C
                                        see include deck elccmt for details
     kerr (int,sc,inout) - error flag up to this point.
```

2.1.3.3.1. Subroutines uex101 through uex105

The source code for subroutines uex101, uex102, uex103, uex104, and uex105 is identical to the code for subroutine uex100 listed above.

2.1.3.4. Subroutine uel100 (Calculating Element Matrices, Load Vectors, and Results)

```
*deck.uel100
                                  USERDISTRIB
     subroutine uel100 (elem,ielc,elmdat,eomaskL,nodes,locsvrL,kelreq,
     x kelfil,nr,xyz,u,kelout,zs,zass,damp,gstif,zsc,zscnr,elvol,elmass,
    x center,elener,edindxL,lcerstL)
c --- general lumped mass is demonstrated -----
c *** primary function:
        1. compute element matrices, load vectors, and results
С
c *** secondary functions:
         2. maintain element solution data
c *** user programmable functions may not be used in parallel processing ***
c *** Notice - This file contains ANSYS Confidential information ***
          *** Copyright ANSYS. All Rights Reserved.
          *** ansys, inc.
 input arguments:
        elem
                                   - element label (number)
             (int,ar(IELCSZ),in) - array of element type characteristics
C
                                     (IELCSZ = array size, defined in echprm)
C
        elmdat (int,ar(EL_DIM),in) - array of element data
C
        eomaskL(LONG,sc,in)
                                   - bit pattern for element output
C
С
                                    (see outpcm)
                                   - array of element node numbers
       nodes (int,ar(nnod),in)
C
                                   (nnod = number of nodes; 1 in this case)
C
        locsvrL(LONG,sc,in)
                                   - location of the saved variables
С
C
                                               on file .esav for this element
        kelreq (int,ar(10),in)
                                   - matrix and load vector form requests
С
С
                                   (indices for kelreq are given with output
                                                            arguments below)
C
С
        kelfil (int,ar(10),in)
                                   - keys indicating incoming matrices and
C
                                     load vectors (indices for kelfil are the
                                     same as given for kelreq with output
C
                                    arguments below)
С
                                   - matrix and load vector size
С
        nr
               (int.sc.in)
               (dp,ar(6,nnod),in) - nodal coordinates (orig) and rotation angle
C
        XVZ
                                   - element nodal solution values
               (dp,ar(nr,5),in)
С
  output arguments:
       kelout (int,ar(10),out)
                                   - keys indicating created matrices and
C
                                          load vectors (indices for kelout
С
                                             are the same as for kelreq below,
С
С
                                             as well as kelin and kelout later)
               (dp,ar(nr,nr),inout)- stiffness/conductivity matrix (kelreq(1))
C
        zs
С
               (dp,ar(nr,nr),inout) - mass matrix
               (dp,ar(nr,nr),inout) - damping/specific heat matrix (kelreq(3))
        damp
```

```
(dp,ar(nr,nr),inout) - stress stiffness matrix (kelreq(4))
С
С
       zsc
              (dp,ar(nr),out) - applied f vector
                                                                (kelreq(5))
                                 - n-r restoring f vector
       zscnr (dp,ar(nr),out)
                                                                (kelreg(6))
C
С
                                  or imaginary f vector
                                                                (kelreq(7))
С
       elvol (dp,sc,out)
                                 - element volume
       elmass (dp,sc,out)
                                 - element mass
С
       center (dp,ar(3),out)
                                 - centroid location
С
                                 - element energies
       elener (dp,ar(5),out)
С
       edindxL(LONG,ar(*),out) - element result data file indexes
C
       lcerstL(LONG,sc,inout)
                                 - position on result file
```

2.1.3.4.1. Subroutines uel101 through uel105

The input and output arguments for subroutines uel101, uel102, uel103, uel104, and uel105are identical to subroutine uel100 listed above.

2.1.3.5. Subroutine uep100 (Printing Output for User Elements in POST1 via PRESOL, ELEM)

```
*deck,uep100
                                USERDISTRIB
     subroutine uep100 (iott,elem,nodes,mat, kept,tem,
    x kemn, fluen, kems, force, kens, sig, keel, epel,
    x keth,eptho,epswel,epino, kenl,sigepl,sigrat,hpres,epeq,
    x kepl,eppl, kecr,epcrp)
С
c *** primary function:
                         produce printed output for user100
С
         *** Copyright ANSYS. All Rights Reserved.
C
         *** ansys, inc.
C
c *** Notice - This file contains ANSYS Confidential information ***
  ******* this subroutine is provided for user information *******
c *** user programmable features may not be used in parallel processing ***
c input arguments:
     iott (int,sc,in)
                                - output unit number
C
С
     elem
              (int,sc,in)
                                 - element number
     nodes
                                - node numbers
С
             (int,ar(2),in)
             (int,sc,in)
                                - material number
C
     mat.
С
    kept
             (int,sc,in)
                                - key to print temperatures
     tem
             (dp,ar(2),in)
С

    nodal temperatures

     kemn
             (inr,sc,in)
                                 - key to print fluences
С
     fluen
              (dp,ar(2),in)
                                 - neutron fluences
С
                                - key to print moment forces
C
     kems
             (int,sc,in)
     force (int,sc,in)
                                - member force fx
С
С
     kens
             (int,sc,in)
                                - key to print strains
     sig
             (dp,sc,in)
                                - axial stress
С
              (int,sc,in)
С
     keel
                                 - key to print elastic strain
                                - axial elastic strain
С
     epel
              (dp,sc,in)
                                - key to print thermal, initial, swelling strai
     ket.h
             (int,sc,in)
С
                                - axial thermal strain
С
     eptho
             (dp,sc,in)
С
     epswel (dp,sc,in)
                                - swelling strain
             (dp,sc,in)
С
     epino
                                - initial axial strain
     kenl
              (int,sc,in)
                                - key set if any nonlinear materials present
С
                                - stress in stress-strain curve
С
     sigepl
              (dp,sc,in)
С
     sigrat
              (dp,sc,in)
                                - stress ratio
С
     hpres
              (dp,sc,in)
                                - hydrostatic pressure
     epeq
С
              (dp,sc,in)
                                - plastic equivalent strain
С
     kepl
              (int,sc,in)
                                 - key to print plastic strain
С
     eppl
              (dp,sc,in)
                                - plastic strain
                                - key to print creep strain
     kecr
              (int,sc,in)
C
С
     epcrp (dp,sc,in)
                                 - creep strain
```

```
c output arguments: none
c
```

2.1.3.5.1. Subroutines uep101 through uep105

The source code for subroutines uep101, uep102, uep103, uep104, and uep105 is identical to subroutine uep100 listed above.

2.1.3.6. Subroutine usertr (Adjusting the Nodal Orientation Matrix)

```
*deck,usertr
                                  USERDISTRIB
     subroutine usertr (node,tr)
c *** primary function: adjust nodal orientation matrix
      secondary function: study nodal orientation matrix
С
        accessed with ielc(notran) = -100
          *** Copyright ANSYS. All Rights Reserved.
          *** ansys, inc.
С
С
c *** Notice - This file contains ANSYS Confidential information ***
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
С
С
c input arguments:
С
     variable (typ,siz,intent)
                                  description
С
     node
               (int,sc,in)
                                 - node number being acted upon
               (dp,ar(32,32),inout) - nodal to global orientation matrix
C
С
c output arguments:
     variable (typ,siz,intent)
                                 description
C
               (dp,ar(32,32),inout) - nodal to global orientation matrix
С
С
           tr is a matrix that is already defined based on the degrees
С
С
           of freedom selected.
          it does not normally need to be changed.
C
С
           it may be printed out here to study. its functional size is
          nr by nr, where nr is the number of degrees of freedom in the
C
           element
С
```

2.1.3.7. Subroutine userac (Accessing Element Information)

This subroutine is provided for demonstration purposes.

```
*deck,userac
                                  USERDISTRIB
     subroutine userac (elem)
c *** primary function: To demonstrate user access of element information.
c --- Given the element number, all information about the element is avaiable.
c --- Starting with elmdat, one can get the element type, real constant number,
c --- the material number, the element coordinate system number, as well as
c --- the node numbers. Then, one can get more information about any or all
c --- of these things. The below demonstrates getting the element type and
c --- real constants.
          *** Copyright ANSYS. All Rights Reserved.
С
          *** ansys, inc.
C
c *** Notice - This file contains ANSYS Confidential information ***
C
С
  input arguments:
С
     variable (typ,siz,intent)
                                  description
                            - element number
     elem (int,sc,in)
```

```
c output arguments:
c none
c
```

2.2. Supporting Subroutines for Element Creation

The following subroutines support the user subroutines for creating a new element (p. 149) (using the database-access method (p. 169)):

- 2.2.1. Subroutine nminfo (Returning Element Reference Names)
- 2.2.2. Subroutine sygidx (Fetching the Index for Saved Variables)
- 2.2.3. Subroutine syrget (Fetching Saved Variable Data for an Element)
- 2.2.4. Subroutine syrput (Writing an Element's Saved Variable Set)
- 2.2.5. Subroutine sypidx (Writing the Saved Variable Element Index to a File)
- 2.2.6. Subroutine mreuse (Determining Which Element Matrices Can Be Reused)
- 2.2.7. Subroutine subrd (Reading Element Load Data for a Substructure Generation Run)
- 2.2.8. Subroutine subwrt (Writing an Element Load Vector to a File for a Substructure Generation Run)
- 2.2.9. Subroutine rvrget (Fetching Real Constants for an Element)
- 2.2.10. Subroutine propev (Evaluating a Group of Material Properties)
- 2.2.11. Subroutine prope1 (Evaluating One Material Property)
- 2.2.12. Subroutine pstev1 (Evaluating Material Properties for 1-D Elements)
- 2.2.13. Subroutine tbuser (Retrieving User Table Data)
- 2.2.14. Subroutine plast1 (Updating an Element's Plastic History)
- 2.2.15. Subroutine plast3 (Updating an Element's Plastic History, 4 or 6 components)
- 2.2.16. Subroutine creep1 (Updating an Element's Creep History)
- 2.2.17. Subroutine creep3 (Updating an Element's Creep History, 3-D Elements)
- 2.2.18. Subroutine swell1 (Updating an Element's Swelling History)
- 2.2.19. Subroutine swell3 (Updating an Element's Swelling History, 3-D Elements)
- 2.2.20. Function elLenPsvrBuf (Determining Additional ESAV Record for Plasticity)
- 2.2.21. Function nlget (Retrieving Material Nonlinear Property Information)
- 2.2.22. Subroutine usereo (Storing Data in the nmisc Record)
- 2.2.23. Subroutine eldwrtL (Writing Element Data to a File)
- 2.2.24. Subroutine eldwrnL (Writing Element Nonsummable Miscellaneous Data to the Results File)
- 2.2.25. Subroutine trrot (Calculating the Rotation Vector)
- 2.2.26. Subroutine rottr (Calculating the Transformation Matrix)
- 2.2.27. Subroutine xyzup3 (Updating an Element's 3-D Nodal Coordinates)
- 2.2.28. Subroutine tmpget (Defining Current Temperature Loads)
- 2.2.29. Subroutine prsget (Defining Current Pressure Loads)
- 2.2.30. Subroutine cnyget (Defining Current Convection Loads)
- 2.2.31. Subroutine hgnget (Defining Current Heat Generation Loads)

2.2.32. Subroutine prinst (Calculating Principal Stress and Stress Intensity)

2.2.1. Subroutine nminfo (Returning Element Reference Names)

```
*deck,nminfo
     subroutine nminfo (ielc,rname)
c *** primary function:
                       set element reference names
c *** secondary functions: none
       ----- to get name back, use nameiq
C
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
    variable (typ,siz,intent)
C
                                description
     ielc (int,ar(*),inout) - element characteristic vector
С
    rname (chr,sc,in)
                              - 8 character reference name
C
С
С
 output arguments:
   variable (typ,siz,intent)
                                description
C
    ielc (int,ar(*),inout) - element characteristic vector with
С
                                  element name encoded
С
```

2.2.2. Subroutine sygidx (Fetching the Index for Saved Variables)

```
*deck,svgidx
     subroutine sygidx (locsvr,svindx)
c *** primary function: get the index for saved variables
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
С
     locsvr (LONGINT,sc,in) - pointer to location of index
  output arguments:
C
     svindx (int,ar(20),out) - the 20 word index of svr variables
C
                                   1,2-starting loc of this eles svr sets
С
                                   3- length of eles svr sets
C
                                   4-11-relative starting loc for each set
С
                                      4-structural svrs
С
                                      5-thermal/electric/fluid svrs
C
С
                                      6-magnetic svrs
                                      7-nonlinear syrs
C
С
                                      8-plasticity svrs
С
                                      9-creep svrs
                                     10-coupled svrs
C
C
                                     11-user svrs
                                     12-initial strain svrs
С
С
                                     13-section data after FiberSIM conversion
                                                                 (shell181 only)
С
С
                                     14-20 spares
```

2.2.3. Subroutine syrget (Fetching Saved Variable Data for an Element)

```
*deck,svrget
subroutine svrget (svindx,nset,nsvr,svr)
c *** primary function: get svr data set for an element
c *** Notice - This file contains ANSYS Confidential information ***
```

```
c input arguments:
     svindx (int,ar(20),in) - index for svr for this element (see svgidx)
C
              (int,sc,in)
                              - the set number in this index
C
С
                                   = 1 - structural svrs
                                   = 2 - thermal/electric/fluid syrs
C
                                      3 - magnetic svrs
С
                                   = 4 - nonlinear svrs
С
                                   = 5 - plasticity svrs
C
                                   = 6 - creep svrs
                                   = 7 - coupled svrs
С
                                     8 - user svrs
С
С
                                   = 9 - initial stress svrs
                                                  (2,42,82,45,92,95 only)
C
                                   = 10 - section data after FiberSIM conversion
С
С
                                                                (shell181 only)
                                   = 11-17 - spares (note that the first three
С
                                             items in svindx are not available)
С
С
     nsvr
              (int,sc,inout)
                                 - number of dp words expected in this set
 output arguments:
    nsvr (int,sc,inout)
                                - number of dp words in this set
C
              (dp,ar(nsvr),in) - data in this set
С
```

2.2.4. Subroutine syrput (Writing an Element's Saved Variable Set)

```
*deck.svrput
     subroutine svrput (svindx,nset,leng,svr)
c *** primary function:
                          write out a svr data set for an element
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
      svindx (int,ar(20),inout) - the index for svr for this element
                                                                     (see svqidx)
C
               (int,sc,in)
                                  - the set number in this index (same as syrget)
С
                                    = 1 - structural svrs
С
                                    = 2 - thermal/electric/fluid svrs
C
                                    = 3 - magnetic svrs
С
С
                                    = 4 - nonlinear svrs
                                    = 5 - plasticity svrs
= 6 - creep svrs
C
С
                                    = 7 - coupled svrs
C
                                    = 8 - user svrs
С
С
                                    = 9 - initial stress svrs
                                                  (2,42,82,45,92,95 only)
C
                                    = 10 - section data after FiberSIM conversion
С
С
                                                                  (shell181 only)
                                    = 11-17 - spares (note that the first three
C
                                              items in svindx are not available)
С
С
      leng
               (int,sc,in)
                                 - number of dp words in this set
С
               (dp,ar(leng),in) - data in this set
  output arguments:
      svindx (int,ar(10,2),inout) - updated index
```

2.2.5. Subroutine sypidx (Writing the Saved Variable Element Index to a File)

```
*deck,svpidx
subroutine svpidx (locsvr,svindx)
c *** primary function: write the svr element index onto file
```

```
c *** secondary functions: update the locsvr pointer to next element
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
                                    - pointer to start of svr for element
     locsvr (LONGINT,sc,inout)
C
               (int,ar(10,2),in)
                                     - index to svr for this element
С
      svindx
                                         low and high parts of 64 bit address
С
С
  output arguments:
             (LONGINT,sc,inout)
      locsvr
                                    - pointer to start of svr for next element
```

2.2.6. Subroutine mreuse (Determining Which Element Matrices Can Be Reused)

```
*deck.mreuse
      subroutine mreuse (kelrqq, kelfil, elem, ielc, kmasrt, knlmg, kconve,
     x kpheno,kprop,nprop,prop,propo,krvro,rvr,rvro,amodo,asymo, kelin)
c *** primary function:
          determine which Matrices can be REUSEd and which must be recomputed
           from iteration to iteration.
С
           Note: a few special elements have some supplementary logic
C
           to adjust these results further. No attempt as been made to
С
           include all such logic in these routines.
С
С
           Second note: this logic is essentially the same as the old
           sfrm logic. Hopefully, further simplifications and enhancements
С
           will be made in the future. (Especially in gap elements and in
С
С
           multilayer elements)
           the whole idea of kpheno, a holdover from the sfrm routines,
C
С
           needs to be looked at and possibly eliminated.
c *** Notice - This file contains ANSYS Confidential information ***
С
  input arguments:
С
     kelrqq
               (int,ar(10),in)
                                   - request keys (needed for this analysis)
C
      kelfil
               (int,ar(10),in)
                                   - keys indicating matrices on the file
С
                                   - element number
C
      elem
               (int,sc,in)
               (int,ar(IELCSZ),in) - array of element type characteristics
С
      ielc
С
      kmasrt
              (int,sc,in)
                                   - does the mass matrix have rotational DOF?
                                       0 - no
                                                 1 - yes(with nlgeom, sfrmln)
C
С
      knlmg
               (int,sc,in)
                                   - nonlinear magnetic curve exists in this
С
                                      element
                                       0 - no
                                                   1 - yes
С
      kconve
               (int,sc,in)
                                   - key indicating existence of convections
С
С
                                      in this element
С
                                       0,1 - no
                                                   2 or more - yes
С
                                      must be input as 'i' if not used, as is
                                      changed in this routine(for analyzer).
C
                                      i = 0 must be used in calling routine
С
                                      if kpheno = 1.
С
                                    - key for type of phenomenon/level of check
C
      kpheno
               (int,sc,in)
                                      0 - structural like old sfrmln,1s,3n,3s,fl
С
                                      1 - thermal like old sfrmlc,1t,2t,3t
С
                                      2 - electrical/magnetic like some of old
С
С
                                      3 - general
                                                     like old sfrmoo
С
      kprop
               (int,sc,in)
                                   - key indicating which material properties
C
                                      in the prop vector that need to be
С
                                      checked (see below)
C
С
               (int,sc,in)
                                   - number of properties
      nprop
С
      prop
               (dp,ar(nprop),in)
                                   - current mat props
               (dp,ar(nprop),inout) - previous material properties
C
      propo
С
      krvro
               (int,sc,in)
             = 0 - real constants are used by this element, and the old
```

```
C
                    values(rvro) have been saved; or the element does not
С
                    use real constants. Any change of real constants
                    causes all matrices to be reformed.
C
             = 1 - real constants are used by this element and the old
C
С
                    values(rvro) have been saved. However, any change
                    of real constants will cause the run to terminate,
C
                    because the results would be too unpredictable.
С
С
                    (e.g. gap elements)
             = 2 - element is nonlinear, so do not bother to check
C
             = 3 - real constants are used by this element, and the old
                    values(rvro) have been saved. However, no checking is
С
С
                    done in this routine because of needed customized logic.
             = 4 - real constants are used by this element, but the old
                    values(rvro) have not been saved because it was
C
                    decided not to use this much storage. therefore, no check
C
С
                    can be made to determine if matrices should be reformed.
                    (e.g. 100 layer elements)
С
             = 5 - real constants are used by this element, but the old
С
С
                    values(rvro) have not been saved because the real
                    constants have no effect on matrix formulation.
C
                    (e.g. acoustic elements)
C
               (dp,ar(*),in)
                                   - current real constants
С
     rvr
               (dp,ar(*),inout)
                                   - previous real constants
С
     rvro
                                   - previous value of mode
               (dp,sc,inout)
С
      amodo
                                   - previous value of isym
C
     asvmo
               (dp,sc,inout)
С
С
  output arguments:
    propo (dp,ar(nprop),inout)- current material properties
C
               (dp,ar(*),inout) - current real constants
(dp,sc,inout) - current value of mode
С
      rvro
               (dp,sc,inout)
С
     amodo
               (dp,sc,inout)
                                  - current value of isym
C
     asvmo
      kelin
               (int,ar(10),out) - keys indicating matrices to form
С
C
```

2.2.7. Subroutine subrd (Reading Element Load Data for a Substructure Generation Run)

```
*deck,subrd
      subroutine subrd (iel, key, nd, vect, ka)
c *** primary function:
                           read element load data from file for substructure
С
                            generation run
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
С
С
     iel
              (int,sc,in)
                                  - element number
      key
               (int,sc,in)
                                  - type of load data
C
                                   = 1 temperature
С
                                   = 2 fluences
C
С
                                   = 3 heat generation rates
                                   = 4 current densities
C
                                   = 9 end pressures (needed for beams/pipes)
C
С
                                   =10 pressures
С
                                   =11 film coefficients
                                   =12 bulk temperatures
С
С
                                   =13 extra displacement shapes
                                   =14 thermal strains(eptho in el42)
С
                                   =15 thermal flux (as in el55)
C
                                   =16 initial strains(epino in el01)
С
                                   =17 magnetic virtual displacements
C
С
                                   =18 calculated source field(hsn in el96)
С
                                   =20 element load vector
                                   =30 copy - do not scale(tempev in el42)
C
                                       first load step only
C
          (int,sc,in)
                                  - number of data items
```

```
c output arguments:
c vect (dp,ar(nd),out) - array of load data
c ka (int,sc,out) - load activation key
c = 0 no load for this data
c = 1 load is active
```

2.2.8. Subroutine subwrt (Writing an Element Load Vector to a File for a Substructure Generation Run)

```
*deck,subwrt
     subroutine subwrt (iel,nvect,kkey,nd,vect,ref)
c *** primary function: write element load vect to file for substructure
                          generation run
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
    iel
            (int,sc,in)
                                - element number
С
С
     nvect
            (int,sc,in)
                                - number of load vectors
                                  (current load step number)
C
     kkey
             (int,sc,in)
                                - type of load vect
С
                                 = 1 temperature
С
                                 = 2 fluences
C
                                 = 3 heat generation rates
                                 = 4 current densities
С
С
                                 = 9 end pressures
С
                                 =10 pressures
                                 =11 film coefficients
C
С
                                 =12 bulk temperatures
С
                                 =13 extra displacement shapes
C
                                 =14 thermal strains(eptho in el42)
С
                                 =15 thermal flux (as in el55)
                                 =16 initial strains(epino in el01)
C
                                 =17 magnetic virtual displacements
С
                                 =18 calculated source field(hsn in el96)
С
                                 =20 element load vector
C
                                 =30 copy - do not scale(tempev in el42)
С
С
     nd
              (int,sc,in)
                               - number of vect items
              (dp,ar(nd),in) - array of load data
     vect.
С
С
    ref
              (dp,sc,in)
                                - reference value for zero load
c output arguments: none
```

2.2.9. Subroutine rvrget (Fetching Real Constants for an Element)

```
*deck,rvrget
     subroutine rvrget (iel,ireal,ielc,nrvr,rvr)
c *** primary function: get the real constants for an element
    typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                              intent=in,out,inout
С
     variable (typ,siz,intent)
                                description
               (int,sc,in) - element number
       iel
С
        ireal
                (int,sc,in)
                                 - real constant set number
C
               (int,ar(*),in)
С
                                 - elment type characteristics
     output arguments:
C
        nrvr (int,sc,out) - number of real variables
```

```
c rvr (dp,ar(*),out) - element real constants
c *** mpg magnetic element usage - iel ?
```

2.2.10. Subroutine propev (Evaluating a Group of Material Properties)

```
*deck,propev
      subroutine propev (iel,mtr,lp,tem,prop,n)
c *** primary function:
                         to evaluate a group of material properties
      propev is used to pass two or more material property numbers
      thru the lp array to determine which temperature dependent
C
      material properties are to be evaluated.
С
      thus, the 3 propel calls:
С
         call propel (elem,mat, 1,tem,e(1))
C
         call propel (elem, mat, 10, tem, alpha)
         call propel (elem, mat, 13, tem, dens)
C
      should be combined as:
С
         integer lp(3)
         data lp /1,10,13/
С
         call propev (elem,mat,lp(1),tem,prop(1),3)
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
     iel (int,sc,in)
                             - element number
С
      mtr (int,sc,in)
                             - material number(input quantity mat, mat comma
C
         (int,ar(n),in)
                             - keys for which specific value is requested
С
                                      each group must be in ascending
C
С
                                      order (ex,ey,ez, etc)
С
                                   if negative, a required property
C
                                   if zero, leave prop term unchanged
С
         ---- MP command labels --
        EX = 1, EY = 2, EZ = 3, NUXY = 4, NUYZ = 5, NUXZ = 6, GXY = 7, GYZ = 8,
С
        GXZ = 9, ALPX=10, ALPY=11, ALPZ=12, DENS=13, MU =14, DAMP=15, KXX =16,
C
        KYY =17, KZZ =18, RSVX=19, RSVY=20, RSVZ=21, C =22, HF =23, VISC=24,
        EMIS=25, ENTH=26, LSST=27, PRXY=28, PRYZ=29, PRXZ=30, MURX=31, MURY=32,
С
        MURZ=33, PERX=34, PERY=35, PERZ=36, MGXX=37, MGYY=38, MGZZ=39, EGXX=40,
С
С
        EGYY=41, EGZZ=42, SBKX=43, SBKY=44, SBKZ=45, SONC=46, DMPS=47, ELIM=48,
        USR1=49, USR2=50, USR3=51, USR4=51, FLUI=53, ORTH=54, CABL=55, RIGI=56,
C
С
        HGLS=57, BVIS=58, QRAT=59, REFT=60, CTEX=61, CTEY=62, CTEZ=63, THSX=64,
С
        THSY=65, THSZ=66, DMPR=67, LSSM=68, BETD=69, ALPD=70, RH =71, DXX =72,
        DYY =73, DZZ =74, BETX=75, BETY=76, BETZ=77, CSAT=78, CREF=79, CVH =80
C
                                               (see mpinit for uncommented code)
C
С
      (see chapter 2 of the elements volume of the user's manual
       for a detailed description))
C
С
      t.em
               (dp,sc,in)
                                 - temperature at which to evaluate material
               (int,sc,in)
                                 - number of properties to be evaluated.
C
                                    (20 maximum)
С
                                    If n = 1, use propel instead.
C
  output arguments:
              (dp,ar(n),out)
                                 - values of material property
     prop
```

2.2.11. Subroutine prope1 (Evaluating One Material Property)

```
*deck,propel
```

```
subroutine propel (iel,mtr,icon,tem,prop1)
c *** primary function: to evaluate one material property
                                                                                        (if multiple material properties are to
C
                                                                                          be evaluated, use propev)
C
c *** secondary functions: to ensure that certain required props are present
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
                iel
                                       (int,sc,in)
                                                                                                      - element number
                                               (int,sc,in)
                                                                                                      - material number
                  mtr
C
С
                                            (int,sc,in)
                                                                                                      - key for which specific value is requested
С
                                                                                                             (negative if property is required)
                            ---- MP command labels -----
С
                           EX = 1, EY = 2, EZ = 3, NUXY = 4, NUYZ = 5, NUXZ = 6, GXY = 7, GYZ = 8,
                            GXZ = 9, ALPX=10, ALPY=11, ALPZ=12, DENS=13, MU = 14, DAMP=15, KXX = 16,
                             \texttt{KYY} = 17 \,, \; \texttt{KZZ} = 18 \,, \; \texttt{RSVX} = 19 \,, \; \; \texttt{RSVY} = 20 \,, \; \; \texttt{RSVZ} = 21 \,, \; \; \texttt{C} \qquad = 22 \,, \; \; \texttt{HF} \qquad = 23 \,, \; \; \texttt{VISC} = 24 \,, \; \; \texttt{C} = 22 \,, \; \; \texttt{C
С
                            EMIS=25, ENTH=26, LSST=27, PRXY=28, PRYZ=29, PRXZ=30, MURX=31, MURY=32,
С
                            MURZ=33, PERX=34, PERY=35, PERZ=36, MGXX=37, MGYY=38, MGZZ=39, EGXX=40,
С
                           EGYY=41, EGZZ=42, SBKX=43, SBKY=44, SBKZ=45, SONC=46, DMPS=47, ELIM=48,
C
                            USR1=49, USR2=50, USR3=51, USR4=51, FLUI=53, ORTH=54, CABL=55, RIGI=56,
                            HGLS=57, BVIS=58, QRAT=59, REFT=60, CTEX=61, CTEY=62, CTEZ=63, THSX=64,
С
                            THSY=65, THSZ=66, DMPR=67, LSSM=68, BETD=69, ALPD=70, RH =71, DXX =72,
С
                            DYY =73, DZZ =74, BETX=75, BETY=76, BETZ=77, CSAT=78, CREF=79, CVH =80
С
                                                                                                                                                    (see mpinit for uncommented code)
C
С
                  tem
                                               (dp,sc,in)
                                                                                                       - temperature at which to evaluate material
С
      output arguments:
                                                                                                       - value of material property
                  prop1
                                           (dp,sc,out)
```

2.2.12. Subroutine pstev1 (Evaluating Material Properties for 1-D Elements)

```
*deck,pstev1
     subroutine pstev1 (elem, matin, tem, prop)
c *** primary function: to evaluate material properites for 1-d elements
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
            (int,sc,in)
                                  - element number (for anserr)
     elem
C
С
     matin
              (int,sc,in)
                                 - material reference number
                                   if negative, no required properties
C
C
     tem
             (dp,sc,in)
                                  - temperature for evaluation
c output arguments:
             (dp,ar(5),out)
С
                                 - material properties: ex,nuxy,gxy,alpx,dens
```

2.2.13. Subroutine tbuser (Retrieving User Table Data)

```
c tbprop (dp,ar(numitm),out) - array of tb data
```

2.2.14. Subroutine plast1 (Updating an Element's Plastic History)

```
*deck,plast1
     subroutine plast1 (option,elem,intpt,mat,kstartL,tem,dtem,e,
                            ktform,dens,flu,dflu,epel,eppl,statev,usvr,
                            epeq,plwork,sigepl,sigrat,et)
c *** primary function:
                           to update the plastic history (for 1 component)
                             used by: LINK1, LINK8, BEAM23, BEAM24, and
С
                                         SOLID65(reinforcing)
c *** secondary functions: to compute the material tangent matrix if requested
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
     option (int,sc,in)
                                     - plasticity option
                                     - element number (label)
      elem (int,sc,in)
С
С
      intpt
                (int,sc,in)
                                      - element integration point number
                                      - material reference number
      mat
                (int,sc,in)
С
     kstartL (intL,sc,in)
                                     - virtual starting address of the data table
С
                                     - temperature at the end of this substep
С
     tem (dp,sc,in)
С
     dtem (dp,sc,in)
                                     - temperature increment over this substep
                (dp,sc,in)
                                      - elastic modulus
C
      e
                                      - request key for tangent matrix formation
      ktform (int,sc,in)
С
     dens (dp,sc,in)
                                     - material density
С
                                     - fluence at the end of this substep
С
     flu (dp,sc,in) - fluence at the end of this substep

dflu (dp,sc,in) - fluence increment over this substep

epel (dp,sc,inout) - modified total strain (trial strain)

eppl (dp,sc,inout) - plastic strain at previous substep

statev (dp,ar(6),inout) - state variables at previous substep
С
С
С
С
                (dp,ar(*),inout) - user-defined state variables (for userpl)
     usvr
С

      (dp,sc,inout)
      - effective plastic strain at previous

      (dp,sc,inout)
      - accumulated plastic work at prev substep

                                     - effective plastic strain at prev substep
С
     epeq
С
     plwork (dp,sc,inout)
c output arguments:
      epel (dp,sc,inout)
                                      - elastic strain
С
      eppl
                                     - updated plastic strain
                (dp.sc.inout)
C
     statev (dp,ar(6),inout) - updated state variables
С
     usvr (dp,ar(*),inout) - updated user-defined state variables
С
     epeq
     epeq (dp,sc,inout) - updated effective plastic strain
plwork (dp,sc,inout) - updated accumulated plastic work
                                      - updated effective plastic strain
С
С
     sigepl (dp,sc,out)
                                     - stress value on stress-strain curve
С
C
     sigrat (dp,sc,out)
                                     - ratio of trial stress to yield stress
     et
               (dp,sc,out)
                                      - tangent modulus
С
  internal variables:
     deppl (dp,sc)
                                       - equivalent plastic strain increment
```

2.2.15. Subroutine plast3 (Updating an Element's Plastic History, 4 or 6 components)

```
c *** secondary functions: to compute the material tangent matrix if requested
c *** Notice - This file contains ANSYS Confidential information ***
 input arguments:
C
     option
               (int,sc,in)
                                    - plasticity option
С
                                   - element number (label)
С
      elem
               (int,sc,in)
                                   - element integration point number
               (int,sc,in)
     int.pt.
C
     mat
               (int,sc,in)
                                   - material reference number
     kstartL (intL,sc,in)
                                   - virtual starting address of the data table
C
С
     ncomp
               (int,sc,in)
                                   - number of stress/strain components (4 or 6)
С
      tem
               (dp,sc,in)
                                   - temperature at the end of this substep
               (dp,sc,in)
                                   - temperature increment over this substep
     dtem
С
                                   - material property array (ex,ey,ez,
С
     prop
               (dp,ar(9),in)
С
                                     gxy,gyz,gxz, uxy,uyz,uxz)
               (dp,ar(ncomp,ncomp),in) - elastic stress-strain matrix
С
               (int,sc,in)
                                   request key for tangent matrix formationmaterial density
     ktform
С
               (dp,sc,in)
С
     dens
     f111
               (dp,sc,in)
                                   - fluence at the end of this substep
C
     dflu
               (dp,sc,in)
                                   - fluence increment over this substep
C
               (dp,ar(ncomp),inout) - modified total strain (trial strain)
     epel
С
     eppl
               (dp,ar(ncomp),inout) - plastic strain at previous substep
С
      statev
               (dp,ar(ncomp,6),inout) - state variables at previous substep
C
               (dp,ar(*),inout) - user-defined state variables (for pluser)
C
     usvr
               (dp,sc,inout)
                                   - effective plastic strain at prev substep
С
     epeq
С
     plwork
               (dp,sc,inout)
                                   - accumulated plastic work at prev substep
C
     kplst
               (int,sc,in)
                                   - plane stress key (form dtt if kplst=1)
  output arguments:
С
     epel
               (dp,ar(ncomp),inout) - elastic strain
C
               (dp,ar(ncomp),inout) - updated plastic strain
C
               (dp,ar(ncomp,6),inout)- updated state variables
C
     statev

    updated user-defined state variables
    updated effective plastic strain

     usvr
               (dp,ar(*),inout)
С
С
     epeq
               (dp,sc,inout)
                                   - updated accumulated plastic work
     plwork
              (dp,sc,inout)
C
С
     sigepl
              (dp,sc,out)
                                   - stress value on stress-strain curve
С
     sigrat
              (dp,sc,out)
                                   - ratio of trial stress to yield stress
     dt.
               (dp,ar(ncomp,ncomp),out) - material modulus modified by dscpar
C
С
     dtt
               (dp,ar(ncomp,ncomp),out)- consistent tangent modulus
                                           (formed only if kplst=1)
С
  internal variables:
      deppl
              (dp,sc)
                                    - equivalent plastic strain increment
```

2.2.16. Subroutine creep1 (Updating an Element's Creep History)

```
*deck,creep1
     subroutine creep1 (option,elem,intpt,mat,kstartL,epel,e,epcrp,
    x statev, usvr, tem, dtem, fluen, dflu, sig)
c *** primary function:
                          to update the creep history for 1-d elements
                           used by: LINK1, LINK8, BEAM23, BEAM24, and
C
                                     SOLID65(reinforcing)
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
C
               (int,sc,in)
С
     option
                                   - creep option
                                   - element number (label)
С
      elem
               (int,sc,in)
     intpt
               (int,sc,in)
                                   - element integration point number
C
                                   - material reference number
С
               (int,sc,in)
     kstartL (intL,sc,in)
                                   - virtual starting address of the data table
C
С
      epel
               (dp,sc,inout)
                                   - elastic strain
               (dp,sc,in)
                                   - elastic modulus
C
                                   - creep strain at previous substep
C
      epcrp
               (dp,sc,inout)
     statev (dp,ar(7),inout) - state variables at previous substep
```

```
C
               (dp,ar(*),inout) - user-defined state variables (for usercr)
С
      tem
               (dp,sc,in)
                           - temperature at the shall - temperature increment over this substep
                                   - temperature at the end of this substep
     dtem
               (dp,sc,in)
C
С
     fluen
               (dp,sc,in)
                                 - fluence at the end of this substep
С
     dflu
               (dp,sc,in)
                                  - fluence increment over this substep
С
     epel
               (dp,sc,inout)
                                   - elastic strain adjusted for creep increment
               (dp,sc,inout)
                                   - stress (not really used)
С
      siq
  output arguments:
C
     epcrp
              (dp,sc,inout)
                                 - updated creep strain
                                - updated state variables
               (dp,ar(7),inout)
С
     statev
С
     usvr
               (dp,ar(*),inout)
                                  - updated user-defined state variables
     sig
               (dp,sc,inout)
                                  - stress (recomputed if requested)
```

2.2.17. Subroutine creep3 (Updating an Element's Creep History, 3-D Elements)

```
*deck,creep3
     subroutine creep3 (option,elem,intpt,mat,kstartL,ncomp,epel,e,
    x posn,d,epcrp,statev,usvr,tem,dtem,fluen,dflu,kplst,sig,hsig)
c *** primary function: to update the creep history for 3-d elements
C
       used by: PLANE13, PIPE20, PLANE42, SHELL43, SOLID45,
                 PIPE60, SOLID62, SOLID65, PLANE82, SHELL91,
С
                 SOLID92, SHELL93, SOLID95, SOLID191
С
c *** Notice - This file contains ANSYS Confidential information ***
 input arguments:
С
                              - creep option
С
     option (int,sc,in)
     elem
              (int.sc.in)
                                - element number (label)
C
С
     intpt
             (int,sc,in)
                                - element integration point number
С
     mat
               (int,sc,in)
                                 - material reference number
                                  - virtual starting address of the data table
С
     kstartL (intL,sc,in)
                                  - number of stress/strain components (4 or 6)
С
     ncomp
              (int,sc,in)
              (dp,ar(ncomp),inout)- elastic strain
С
     epel
              (dp,sc,in) - elastic young'S MODULUS
С
              (dp,sc,in)
                                  - poisson'S RATIO
С
    posn
              (dp,ar(ncomp,ncomp),in) - elastic stress-strain matrix
С
     Ы
С
     epcrp
              (dp,ar(ncomp),inout) - creep strain at previous substep
С
     statev
              (dp,ar(ncomp*5+2),inout)- state variables at previous substep
              (dp,ar(*),inout) - user-defined state variables (for usercr)
     usvr
C
              (dp,sc,in)
С
     tem
                                - temperature at the end of this substep
С
     dtem
              (dp,sc,in)
                                - temperature increment over this substep
     fluen
              (dp,sc,in)
                                  - fluence at the end of this substep
С
     dflu
              (dp,sc,in)
                                 - fluence increment over this substep
С
                                 - plane stress/plane strain key
С
     kplst
              (int,sc,in)
С
     siq
              (dp,ar(ncomp),inout) - stresses (not used in input)
     hsig
              (dp,ar(1),inout) - hydrostatic stress (not used in input)
С
С
  output arguments:
              (dp,ar(ncomp),inout) - elastic strain adjusted for creep increment
C
     epel
              (dp,ar(ncomp),inout) - updated creep strain
C
     epcrp
             (dp,ar(ncomp*5+2),inout)- updated state variables
C
С
     usvr
              (dp,ar(*),inout)
                                - updated user-defined state variables
               (dp,ar(ncomp),inout) - stresses (redefined if c13 > 0)
С
     siq
     hsiq
              (dp,sc,inout)
                                  - hydrostatic stress (redefined if c13 > 0)
```

2.2.18. Subroutine swell1 (Updating an Element's Swelling History)

```
*deck,swell1
```

```
subroutine swell1 (option,elem,intpt,mat,kstartL,epswel,epel,e,
     x fluen, dfluen, tem, dtem, usvr)
c *** primary function:
                          to update the swelling history for 1-d elements
С
                           used by: LINK1, LINK8, BEAM23, and BEAM24
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
             (int,sc,in)
                                  - swelling option
     option
C
      elem
               (int,sc,in)
                                  - element number (label)
     intpt
               (int,sc,in)
                                  - element integration point number
C
                                   - material reference number
С
     mat.
               (int,sc,in)
С
     kstartL (intL,sc,in)
                                  - virtual starting address of the data table
               (dp,sc,inout)
                                  - swell strain at previous substep
     epswel
С
               (dp,sc,inout)
                                   - elastic strain
С
     epel
С
               (dp,sc,in)
                                   - elastic young'S MODULUS
     fluen
               (dp,sc,in)
                                   - fluence at the end of this substep
С
     dfluen
               (dp,sc,in)
                                   - fluence increment over this substep
С
С
     t.em
               (dp,sc,in)
                                  - temperature at the end of this substep
     dtem
               (dp,sc,in)
                                  - temperature increment over this substep
C
               (dp,ar(*),inout)

    user-defined state variables (for usersw)

C
     usvr
С
  output arguments:
               (dp,sc,inout)
                                   - elastic strain adjusted for swelling inc
C
                                   - updated swelling strain
C
      epswel
               (dp.sc.inout)
                                   - updated user-defined state variables
               (dp,ar(*),inout)
```

2.2.19. Subroutine swell3 (Updating an Element's Swelling History, 3-D Elements)

```
*deck,swell3
     subroutine swell3 (option,elem,intpt,mat,kstartL,ncomp,epswel,
                         epel, e, nuxy, fluen, dfluen, tem, dtem, usvr)
c *** primary function:
                           to update the swelling history for 3-d elements
        used by: PLANE13, PIPE20, PLANE42, SHELL43, SOLID45,
С
                  PIPE60, SOLID62, PLANE82, SHELL91, SOLID92,
С
                  SHELL93, SOLID95, SOLID191
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
С
     option
              (int,sc,in)
                                   - swelling option
C
С
      elem
               (int,sc,in)

    element number (label)

      intpt
               (int,sc,in)
                                   - element integration point number
С
      mat
               (int,sc,in)
                                   - material reference number
С
                                   - virtual starting address of the data table
С
     kstartL (intL,sc,in)
     ncomp
               (int,sc,in)
                                  - number of stress/strain components (4 or 6)
C
      epswel
               (dp,sc,inout)
                                  - swell strain at previous substep
С
               (dp,ar(ncomp),inout) - elastic strain
      epel
C
                                  - elastic young'S MODULUS
С
               (dp,sc,in)
                                   - poisson'S RATIO
С
      nuxy
               (dp,sc,in)
                                  - fluence at the end of this substep
               (dp,sc,in)
C
      fluen
      dfluen
               (dp,sc,in)
                                   - fluence increment over this substep
С
С
      tem
               (dp,sc,in)
                                   - temperature at the end of this substep
                                   - temperature increment over this substep
      dtem
               (dp,sc,in)
С
С
      usvr
               (dp,ar(*),inout)
                                   - user-defined state variables (for usersw)
  output arguments:
C
      epel
               (dp,ar(ncomp),inout) - elastic strain adjusted for swelling inc
                                 - updated swelling strain
      epswel
               (dp,sc,inout)
C
C
      usvr
               (dp,ar(*),inout)
                                   - updated user-defined state variables
```

2.2.20. Function elLenPsvrBuf (Determining Additional ESAV Record for Plasticity)

```
*deck,elLenPsvrBuf
    function elLenPsvrBuf (mat, plOpt, ncomp)
*** primary function:
          determine additional esave record for plasticity
C
С
   input arguments
   =========
C
          (int,sc,in)
                        - material ID
С
         (int,sc,in)
   plOpt
                       plasticity optionnumber of strain components (1,4, or 6)
C
С
   ncomp (int,sc,in)
   output arguments
C
С
    elLenPsvrBuf (int,sc,out) - number of extra data items saved
С
    local variables
С
    ==========
```

2.2.21. Function nlget (Retrieving Material Nonlinear Property Information)

```
*deck,nlget
     function nlget (mat,iprop,prop)
c *** primary function: get a material non-linear property (TB) table.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
      variable (typ,siz,intent) description
C
        mat (int,sc,in) - material number iprop (int,sc,in) - property number (tbpnum in tblecm)
С
С
                                       use 13 for tb,user
C
                                        use 14 for tb,nl
С
C
    output arguments:
      variable (typ,siz,intent) description
       nlget (int,sc,out) - number of property values
C
С
       prop (dp,ar(nlget),out) - vector of the property values
                                      (the first 15(tbhdsz) items are a header,
                                      given below. The terms are defined in
C
С
        --- terms of the descriptor record:
C
        header(1) = tbtyp
C
        header(2) = tbtems
С
        header(3) = temloc
С
       header(4) = dprtem
C
C
      header(5) = tbrow
C
      header(6) = tbcol
        header(7) = rowkey
C
С
        header(8) = nxtloc
      header(9) = nxttem
C
С
       header(10) = temptr
C
        header(11) = tbpt
        header(12) = tbsiz
C
С
        header(13) = tbopt
С
        header(14) = hypopt
```

```
c header(15) = tbnpts
```

2.2.22. Subroutine usereo (Storing Data in the nmisc Record)

```
*deck, usereo
     subroutine usereo (elem,iout,nbsvr,bsvr,nnrsvr,nrsvr,npsvr,psvr,
    x ncsvr,csvr,nusvr,usvr,nnode,nodes,xyz,vol,leng,time,
    x timinc,nutot,utot,maxdat,numdat,udbdat)
c *** primary function: to call userou, which allows user to store
                               data in nmisc record
c *** Notice - This file contains ANSYS Confidential information ***
С
c input arguments:
   variable (typ,siz,intent)
                                  description
C
    elem (int,sc,in)
                                  - element number
   iout
            (int,sc,in)
                                  - output unit number
С
С
    nbsvr
            (int,sc,in)
                                  - number of basic element variables
                                 - basic element variables
    bsvr
            (dp,ar(nbsvr),in)
С
   nnrsvr (int,sc,in)
                                  - number of nonlinear element variables
С
  nrsvr (dp,ar(nnrsvr),in) - nonlinear element variables
С
С
  npsvr (int,sc,in)
                                 - number of plasticity element variables
            (dp,ar(npsvr),in)
                                  - plasticity element variables
   psvr
C
                                 - number of creep element variables
С
    ncsvr (int,sc,in)
            (dp,ar(ncsvr),in)
С
    csvr
                                 - creep element variables
    nusvr (int,sc,in)
                                - number of user-supplied element variables
С
С
   usvr
            (dp,ar(nusvr),in)
                                - user-supplied element variables
    nnode (int,sc,in)
                                  - number of nodes
С
    nodes (int,ar(nnode),in)
                                  - node numbers
С
            (dp,ar(6,nnode),in) - nodal coordinates and rotations (virgin)
С
    XYZ

    (dp,sc,in)
    - element volume (of all)

    (dp,sc,in)
    - element length (beams,spars,etc)

    vol
С
С
   leng
   time
С
            (dp,sc,in)
                                 - current time
  timinc (dp,sc,in)
С
                                  - current sub step time increment
    nutot (int,sc,in)
utot (dp,ar(nutot
С
                                  - length of dof solution vector utot
            (dp,ar(nutot),in)
C
                                  - solution vector
    maxdat (int,sc,in)
                                  - size of user output array (3 x nnode)
C
                                    actually, = ielc(nmnmup)
С
                                     for contact element it is equale to nusvr
                                     but it does not exceed 120
С
c output arguments:
С
  variable (typ,siz,intent)
                                 description
  numdat (int,sc,out)
                                  - number of user output items in array udbdat
    udbdat (dp,ar(maxdat),out) - user output items to be placed at the end
C
                                     of the nmisc record
С
C
```

2.2.23. Subroutine eldwrtL (Writing Element Data to a File)

```
*deck,eldwrtL
      subroutine eldwrtL (ielem,edtype,lcerstL,edindxL,nval,value)
c *** primary function:
                          output element data to result file
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
С
        ielem
                (int,sc,in)
                                     - element number
        edtype
                                     - element data type (see elparm)
С
                   (int,sc,in)
        lcerstL (LONG,sc,inout) - pointer to results file position
```

```
C
         edindxL (LONG, ar(NUMELEDATASETS), inout) - index to results file data
С
        nval
                   (int,sc,in)
                                     - the total number of values
                                       if edtype = EDEMS,
C
                                           this should -always- be ielc(nmsmis),
C
С
                                           unless there is a variable number, as
                                          in the layered shell elements.
C
        value
                  (dp,ar(nval),in) - output values (real)
```

2.2.24. Subroutine eldwrnL (Writing Element Nonsummable Miscellaneous Data to the Results File)

```
*deck,eldwrnL
     subroutine eldwrnL (elem,ielc,lcerstL,edindxL,nudb,udbdat,
                       nval, value, ndval)
c *** primary function: output element nonsummable miscellaneous data
                                   to result file
С
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
       elem (int,sc,in)
                                - element number
С
С
        ielc (int,ar(IELCSZ),in) - element characteristic vector
                                       defined in elccmt
C
       lcerstL (LONG,sc,inout) - pointer to results file position
С
       edindxL (LONG,ar(NUMELEDATASETS),inout) - index to results file data
С
      nudb
                 (in,sc,inout) - size of what the user wants to add
С
      udbdat (dp,ar(*),in)
                                  - what the user wants to add
C
      nval (int,sc,in)
                                  - the total number of values to
С
                                       be output (does not include nudb)
С
                                       this should -always- be ielc(NMNMIS),
                                       unless there is a variable number, as
C
С
                                      in the layered shell elements.
        value (dp,ar(ndval),in) - output values
С
С
        ndval
                 (int,sc,in) - dimension of value - must be no less than
                                       ielc(NMNMIS) + ielc(NMNMUP)
С
c *** mpg eldwrnL < ell17,el126,el109,el53,el96,el97: write nmisc db \,
```

2.2.25. Subroutine trrot (Calculating the Rotation Vector)

2.2.26. Subroutine rottr (Calculating the Transformation Matrix)

```
*deck,rottr
subroutine rottr (rot,tr)
c primary function: compute transformation matrix from rotation vector ****
```

```
c *** Notice - This file contains ANSYS Confidential information ***
     ref(old): eqn. 20(transposed), rankin and brogan, jpvt,108(1986)165-174.
C
     ref(new): eqn. (b.4), simo and vu-quoc, cmame, 58 (1986), 79-116
С
С
                  (removes singularities at pi and 2*pi)
C
  input arguments:
С
С
     variable (typ,siz,intent) description
                             - rotation parameter in radians
     rot (dp,ar(4),in)
C
c output arguments:
С
     variable (typ,siz,intent)
                                description
              (dp,ar(3,3),out) - transformation matrix corresponding to rot
```

2.2.27. Subroutine xyzup3 (Updating an Element's 3-D Nodal Coordinates)

```
*deck,xyzup3
     subroutine xyzup3 (nnod,u,nr,xyz,nx,xyzup)
c *** primary function: update a 3-d ele nodal coords for large deformation
c *** Notice - This file contains ANSYS Confidential information ***
С
c input arguments:
                                  - number of nodes
     nnod (int,sc,in)
C
              (dp,ar(nr),in)
С
     u
                                  - displacement vector
     nr
              (int,sc,in)
                                  - size of the u vector
C
    xyz
             (dp,ar(nx,nnod),in) - coordinates to be updated
C
С
             (int,sc,in)
                                 - row size of xy
С
c output arguments:
            (dp,ar(3,nnod),out) - updated coordinates
С
C
```

2.2.28. Subroutine tmpget (Defining Current Temperature Loads)

```
*deck, tmpget
     subroutine tmpget (iel,ielc,nnod,nodes,ref,ndat0,begdat,dat,
     x enddat,tlvf)
     primary function: define the current temperature loads
С
c *** Notice - This file contains ANSYS Confidential information ***
С
c typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
c input arguments:
c variable (typ,siz,intent) description
                                - element number
С
    iel (int,sc,in)
    ielc (int,ar(IELCSZ),in) - array of element type characteristics
С
     nnod (int,sc,in) - number of nodes in the nodes array
nodes (int,ar(nnod),in) - list of nodes
С
С
    ref (dp,sc,in) - reference temperature ndat (int,sc,in) - number of data items to get
                                 - reference temperature
С
С
    begdat (dp,ar(ndat),in) - data at the beginning of this load step
C
C
c output arguments:
С
    dat (dp,ar(ndat),out)
                                - data at this time point
     enddat (dp,ar(ndat),out) - data at end of this load step
C
                                 - thermal load vector flag
     tlvf (int,sc,out)
С
С
                                    Should the thermal load vector be computed
С
                                    = 0 - no, temperatures match tref
                                    = 1 - yes, temperatures do not match tref
С
                                    =<0 - no and using table
C
```

```
C Note, that even if tlvf = 0, temperatures may be used to compute temperature-dependent material properties.
```

2.2.29. Subroutine prsget (Defining Current Pressure Loads)

```
*deck,prsget
      subroutine prsget (iel,ielc,nfac,ndat,begdat,dat,enddat,iexist)
     primary function: define the current pressure loads
      See also: PrsRIGet
C
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
        iel
                 (int.sc.in)
                                  - element number
C
С
        ielc (int,ar(IELCSZ),in) - array of element type characteristics
        nfac
                 (int,sc,in) - number of pressure faces
С
                                  - number of pressure values
С
        ndat.
                  (int,sc,in)
        begdat
                (dp,ar(ndat),in) - pressure at the beginning of load step
С
С
     output arguments:
С
        dat
                 (dp,ar(ndat),out) - pressures at this iteration
         enddat
                 (dp,ar(ndat),out) - pressure at end of this load step
C
С
                (int,sc,out)
                                   - flag if pressure exist
                                      = 0 : no pressure
С
                                      = 1 : yes pressure
C
                                      = -1 : no pressure and has table
```

2.2.30. Subroutine cnvget (Defining Current Convection Loads)

```
*deck,cnvget
      subroutine cnvget (iel,ielc,nr,u,nfac,ndat,beghc,begtb,
     x hc,tb,endhc,endtb,iexist)
     primary function: define the current convection loads
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
C
        iel
                  (int,sc,in)
                                   - element number
         ielc (int,ar(IELCSZ),in) - array of element type characteristics
С
                  (int,sc,in) - dimension of u (temperature) vector
(dp,ar(nr),in) - most current temperatures
         nr
C
С
                  (int,sc,in) - number of convection values
С
        nfac
        ndat
C
                  (dp,ar(ndat),in) - hcoef at the beginning of load step
С
         beghc
С
         begtb (dp,ar(ndat),in) - tbulk at the beginning of load step
С
      output arguments:
                  (dp,ar(ndat),out) - hcoef at this substep
С
        hc
С
         t.b
                  (dp,ar(ndat),out) - tbulk at this substep
С
         endhc
                  (dp,ar(ndat),in) - hcoef at the end of this load step
                  (dp,ar(ndat),in) - tbulk at the end of this load step
C
         endtb
         iexist
                                     - flag if convection exist
С
                  (int,sc,out)
С
                                       = 0 - no convection
                                       = 1 - constant convection (with time)
C
                                              does not require new element matrix
С
С
                                       = 2 - changing convection (with time)
С
                                              or deleted convection
                                              requires new element matrix
C
```

2.2.31. Subroutine hgnget (Defining Current Heat Generation Loads)

```
*deck,hgnget
     subroutine hgnget (iel,ielc,nnod,nodes,ndat,begdat,dat,enddat,
     primary function: define the current heat generation loads
С
C
c *** Notice - This file contains ANSYS Confidential information ***
C
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                  intent=in,out,inout
С
С
     input arguments:
C
     variable (typ,siz,intent) description
С
С
       iel
               (int,sc,in)
                                   - element number
        ielc (int,ar(IELCSZ),in) - array of element type characteristics
C
        nnod (int,sc,in) - number of nodes in the nodes array
       nodes (int,ar(nnod),in) - list of nodes
C
        ndat (int,sc,in) - number of data items to get
begdat (dp,ar(ndat),in) - data at the beginning of this load step
С
С
С
    output arguments:
С
С
      dat (dp,ar(ndat),out) - data at this time point
        enddat (dp,ar(ndat),out) - data at end of this load step
C
        iexist (int,sc,out)
                                   - flag if heat generation exist
С
С
                                     = 0 - no heat generation
                                      = 1 - yes heat generation
C
```

2.2.32. Subroutine prinst (Calculating Principal Stress and Stress Intensity)

```
*deck,prinst
     subroutine prinst (s)
    primary function: computes principal stresses and stress intensity
    secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
C
С
     variable (typ,siz,intent)
                                description
              (dp,ar(11),inout) - stress vector
С
C
                  s(1)=sx
С
                  s(2)=sy
                  s(3)=sz
C
С
                  s(4) = sigxy
С
                  s(5) = sigyz
                  s(6) = sigzx
C
С
  output arguments:
C
C
   variable (typ,siz,intent)
                                description
              (dp,ar(11),inout) - stress vector
С
     S
С
                  s(7)=sig1
С
                  s(8) = sig2
C
                  s(9)=sig3
                  s(10)=s.i.
С
С
                  s(11)=sige
c ******* note: all changes to this routine must be made in
С
                 post1 (paprst)
С
```

2.3. Subroutines for Modifying and Monitoring Existing Elements

Following are the user subroutines for modifying or monitoring existing elements:

- 2.3.1. Subroutine userou (Storing User-Provided Element Output)
- 2.3.2. Subroutine useran (Modifying Orientation of Material Properties)
- 2.3.3. Subroutine userrc (Performing User Operations on COMBIN37 Parameters)
- 2.3.4. Subroutine UEIMatx (Accessing Element Matrices and Load Vectors)
- 2.3.5. Subroutine uthick (Getting User-Defined Initial Thickness)
- 2.3.6. Subroutine UsrFictive (Providing User-Defined Fictive Temperature Relationship)
- 2.3.7. Subroutine uflex (Calculating Flexibility Factors for PIPE288 and PIPE289)
- 2.3.8. Subroutine UsrShift (Calculating Pseudotime Time Increment)
- 2.3.9. Subroutine UTimeInc (Overriding the Program-Determined Time Step)
- 2.3.10. Subroutine UCnvrg (Overriding the Program-Determined Convergence)

2.3.1. Subroutine userou (Storing User-Provided Element Output)

```
*deck,userou
                                 USERDISTRIB
     subroutine userou (elem,iout,nbsvr,bsvr,nnrsvr,nrsvr,npsvr,psvr,
     x ncsvr,csvr,nusvr,usvr,nnode,nodes,xyz,vol,leng,time,
    x timinc,nutot,utot,maxdat,numdat,udbdat)
c *** primary function: store user supplied element output
                           in nmisc record
С
С
         in order to activate this user programmable feature,
         the user must enter the usrcal command.
C
С
         *** Copyright ANSYS. All Rights Reserved.
С
         *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
C
С
             this routine is called by almost every element
             the data is stored on the nmisc record.
C
С
             warning: other data may be stored between the
                       documented data and this data.
             in order to see the actual information on the nmisc
C
              record, insert the command:
С
                  dblist,elp,elnum1,elnum2,elinc,11
С
                       where elnum1 = the first element
C
                             elnum2 = the last element
C
                             elinc = the element increment number
C
              after a set command in post1.
C
c input arguments:
   variable (typ,siz,intent)
                                 description
- element number
С
   elem (int,sc,in)
iout (int,sc,in)
С
                                 - output unit number
C
C
  nbsvr (int,sc,in)
                                 - number of basic element variables
  bsvr (dp,ar(nbsvr),in) - basic element variables
С
    nnrsvr (int,sc,in)
                                  - number of nonlinear element variables
C
            (dp,ar(nnrsvr),in)
                                  - nonlinear element variables
С
    nrsvr
    npsvr (int,sc,in)
                                 - number of plasticity element variables
С
    psvr
            (dp,ar(npsvr),in) - plasticity element variables
С
    ncsvr (int,sc,in)
                                 - number of creep element variables
С
    csvr (dp,ar(ncsvr),in) - creep element variables
nusvr (int,sc,in) - number of user-supplied
C
С
                                  - number of user-supplied element variables
                                         (= nstv on the nsvr command)
C
  usvr (dp,ar(nusvr),in) - user-supplied element variables
```

```
- number of nodes
C
    nnode
            (int,sc,in)
            (int,ar(nnode),in)
     nodes
                                  - node numbers
С
            (dp,ar(6,nnode),in) - nodal coordinates and rotations (virgin)
    xyz
C
                                  - element volume (or area if 2-d)
     vol
            (dp,sc,in)
C
С
    leng
            (dp,sc,in)
                                  - element length (beams, spars, etc)
                                  - current time
    time
             (dp,sc,in)
С
     timinc (dp,sc,in)
                                  - current sub step time increment
С
                                  - length of dof solution vector utot
С
    nutot
            (int,sc,in)
            (dp,ar(nutot),in)
                                  - solution vector
C
    ut.ot.
    maxdat (int,sc,in)
                                   - size of user output array (3 x nnode)
                                      for contact element it is equale to nusvr
С
                                      but it dode not exceed 120
С
С
  output arguments:
С
                                   description
   variable (typ,siz,intent)
C
                                   - number of user output items in array udbdat
С
    numdat (int,sc,out)
                                       (maximum size of numdat is ielc(NMNMUP)
С
                                       which is usually three times the number
С
С
                                       of nodes.
                                       For contact elements CONTA171-178, it
C
                                       should be equal or less than NSTV
C
                                       on nsvr command). It cannot exceed 120.
C
     udbdat (dp,ar(maxdat),out)
                                   - user output items to be placed at the end
С
                                     of the nmisc record
C
```

2.3.2. Subroutine useran (Modifying Orientation of Material Properties)

```
*deck,useran
                                  USERDISTRIB
      subroutine useran (vn, vref, elem, thick, xyzctr, bsangl)
      user written routine to modify orientation of material properties
C
      and stresses *****************
       applicable to: shell43,63,91,93,99, solid46,64,191
C
С
        accessed by keyopt
С
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
С
c *** Notice - This file contains ANSYS Confidential information ***
C
c **** warning *** do not change any arguments other than bsangl.
С
                   if you do, your results are probably wrong.
C
C
  input(do not change) ---
            = vector normal to element
C
C
              = unit vector orienting element, essentially edge i-j
              = element number
С
     thick
              = total thickness of element at this point (see note below)
C
     xyzctr = location of element centroid or integration point
С
С
C
     bsangl = output from this subroutine. it represents the angle(s)
C
С
              between vref and the desired orientation. it may have
               the default orientation coming in to useran.
С
                This will be combined with the angles derived from
                the ESYS command.
C
           use 1 angle for 2-d elements and shells
С
           use 3 angles for 3-d solids
С
```

2.3.3. Subroutine userrc (Performing User Operations on COMBIN37 Parameters)

```
USERDISTRIB
*deck,userrc
      subroutine userrc (elem, ireal, type, nusvr, usvr, parm, parmld,
     x c1,c2,c3,c4,fcon)
C
     primary function: user operation on parameter for combin37
      accessed with keyopt(9) = 1
С
          *** Copyright ANSYS. All Rights Reserved.
C
C
         *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
  variable (typ,siz,intent)
                                 description
С
С
    elem
            (int,sc,in)
                                  - element number
С
    ireal
             (int,sc,in)
                                 - element real constant number
                                 - element type number
    type
            (int,sc,in)
C
                                 - number of user-supplied element variables
С
   nusvr (int,sc,in)
С
                                    (input with the NSVR command)
            (dp,ar(nusvr),inout) - user-supplied element variables
С
    usvr
            (dp,sc,in)
                                  - current value of the paramater
С
                                 - value of the parameter at previous time ste
    parmld (dp,sc,in)
С
                                - real constant c1
            (dp,sc,in)
    c1
C
   c2
С
            (dp,sc,in)
                                 - real constant c2
                                  - real constant c3
    c3
            (dp,sc,in)
С
С
            (dp,sc,in)
                                  - real constant c4
C
  output arguments:
С
  variable (typ,siz,intent)
                                 description
C
   usvr (dp,ar(nusvr),inout) - user-supplied element variables
С
                                    may be sent .rst file with usereo
C
                                  - result of calculation
С
    fcon
С
     either c1 or c3 must be nonzero for this logic to be accessed,
C
```

2.3.4. Subroutine UEIMatx (Accessing Element Matrices and Load Vectors)

```
*deck,UElMatx
                                    USERDISTRIB
     subroutine UElMatx (elem,nr,ls,zs,zsc,uelm,ielc,nodes,
                          ElDofEachNode,elmdat,xyzang,lenu)
c primary function:
                       User routine to access element matrices and load vectors.
                       Needs to have USRCAL, UELMATX to be accessed.
С
                       Called after the call to the element routine and
С
                        before the solver.
                       May be used to monitor and/or modify the element matrices
C
C
                                                          and load vectors.
          *** Copyright ANSYS. All Rights Reserved.
C
С
          *** ansys, inc.
      typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
C
С
  input arguments:
     variable (typ,siz,intent)
                                       description
C
               (int,sc,in)
                                    - User element number
С
      elem
               (int,sc,in)
                                    - number of rows in element matrix
     nr
C
С
               (int,ar(nr),in)
                                   - Dof Index vector for this element matrix
С
     ZS
               (dp,ar(nr,nr,*),inout)- K,M,C,SS,KCPLX matrices for this element
               (dp,ar(nr,2),inout) - Element load vector and N-R correction vec
C
      ZSC
               (dp,ar(mr,5),in) - Nodal displacements ---
(int.ar(*),in) - Element type characteristics
      uelm
                                    - Nodal displacements for this element
C
     ielc (int,ar(*),in)
С
```

```
nodes (int,ar(*),in) - Nodes for this element
С
     ElDofEachNode (int,ar(nr),in) - list of dofs for each node in Global
С
     elmdat (int,ar(10),in) - Element data for this element
C
                                  - X,Y,Z,THXY,THYZ,THZX for each element node
     xyzang (dp,ar(6,*),in)
C
C
     lenu
             (int,sc,in)
                                  - Length of global displacement vector
  output arguments:
С
             (dp,ar(nr,nr,4),inout)- K,M,C,SS matrices for this element
С
             (dp,ar(nr,2),inout) - Element load vector and N-R correction vec
C
     WARNING: any CHANGES to these (or any other) arguments will have a direc
      impact on the solution, possibly giving meaningless results. The normal
С
      usage of this routine is simply monitor what is happening.
```

2.3.5. Subroutine uthick (Getting User-Defined Initial Thickness)

```
*deck,uthick
                              USERDISTRIB
    SUBROUTINE uthick (elemId, elemType, matId, realId,
                     numDomIntPts, curCoords, thickness)
*** primary function: get the user defined thickness
С
         *** Copyright ANSYS. All Rights Reserved.
C
С
        *** ansys, inc.
С
    input arguments
С
C
    =========
    Variable (type,sz,i/o) description
С
    elemId
    elemId (int,sc,i) element number
elemType (int,sc,i) element TYPE (181 etc.)
matId (int,sc,i) material number
realId (int,sc,i) real constant set number
С
С
С
С
    numDomIntPts (int,sc,i) number of integration points
С
     curCoords (dp,ar(3,numDomIntPts),i)
С
С
                                current coordinates
С
С
    output arguments
    _____
С
                  (dp,ar(3,numDomIntPts),o)
С
     thickness
                                thickness at the integration points
С
С
С
c --- parameters
```

2.3.6. Subroutine UsrFictive (Providing User-Defined Fictive Temperature Relationship)

```
*deck.UsrFictive
                                      USERDISTRIB
     subroutine UsrFictive (tref,toffst,tem,ftl, veinpt, ftc)
c *** primary function: allow users to write their own
С
                         fictive temperature relationship
                       this logic is accessed with c5 = 11 on the tb, evisc table
c *** secondary function:
                            demonstrate the use of a user-written
                             fictive temperature relationship
C
С
                       this routine could also be used to modify the viscoelastic
С
                       data during solution, i.e., to make the viscoelastic
                       coefficients themselves time-dependent.
C
```

```
c *** notice- this routine contains ansys, inc. confidential information ***
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
С
С
c input arguments:
     variable (type,sze,intent) description
С
              (dp.sc.in) - reference temperature

(dp.sc.in) - temperature offset from absolute zero
С
     tref
     toffst (dp,sc,in)
C
               (dp,sc,in) - temperature at the end of this substep
(dp,sc,in) - previous fictive temperature
С
     ftl
                                    - previous fictive temperature
С
     veinpt (dp,ar(95),inout) - table from tb,evisc
С
С
  output arguments:
С
    variable (type, sze, intent) description
C
С
     veinpt (dp,ar(95),inout) - table from tb,evisc
              (dp,sc,in)
                                    - fictive temperature
С
     ftc
С
```

2.3.7. Subroutine uflex (Calculating Flexibility Factors for PIPE288 and PIPE289)

```
*deck.uflex
                                  USERDISTRIB
     subroutine uflex (elemId,pressInt,pressExt,ex,pois, sflex,twten)
c *** primary function: to (re)compute the flexibility factors
                            for pipe288 and pipe289
С
                           this is accessed by inputting the axial flexibility factor
С
                            as -10.
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
С
          *** Copyright ANSYS. All Rights Reserved.
С
C
          *** ansys, inc.
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
С
C
С
     input arguments:
c elemId
              (int,sc,in) - element number
  pressInt (dp,ar(2),in) - internal pressures at end nodes
pressExt (dp,ar(2),in) - external pressures at end nodes
С
                                        Pressures include hydrostatatic but
С
                                         not hydrodynamic effects.
                (dp,sc,in) - Young's Modulus
c ex
c pois
                (dp,sc,in) - Poisson's ratio
   sflex (dp,ar(6),inout) - input flexibility factors
С
                                (axial, bending about element z,
С
                                 bending about element y, twist, y shear, z shear)
C
c twten
             (dp,sc,inout) - twist-tension factor
C
С
      output arguments:
   sflex (dp,ar(6),inout) - output flexibility factors
С
                                (axial, bending about element z,
C
                                 bending about element y, twist, y shear, z shear)
С
c twten
             (dp,sc,inout) - twist-tension factor
С
```

2.3.8. Subroutine UsrShift (Calculating Pseudotime Time Increment)

```
*deck,UsrShift USERDISTRIB
c Copyright ANSYS. All Rights Reserved.
subroutine UsrShift(dxi,dxihalf,timinc,
```

```
& temp,dtemp,toffst,propsh,nTerms)
С
     calculate pseudotime time increment according
С
     to a user specified shift function
С
c *** Notice - This file contains ANSYS Confidential information ***
С
С
  input arguments:
    timinc (dp,sc,in)
                           - time increment
C
    temp (dp,sc,in)
dtemp (dp,sc,in)
С
                            - current temperature, t_n+1
                            - temperature increment, t_n+1 - t_n
С
    toffst (dp,sc,in)
С
                            - temperature offset to absolute zero
С
                              (specified by TOFFST command)
    propsh (dp,ar,in)
                            - Constants for shift function
С
C
                             (User's input using TB, SHIFT, , , , USER)
С
    nTerms (int,ar,in)
                            - number of user defined constants
                              (specified in TB, SHIFT, , , nTerms, USER)
С
С
  output arguments:
С
   dxi (dp,sc,out)
                            - pseudotime increment
    dxihalf (dp,sc,out)
                            - pseudotime increment over the upper half span
C
```

2.3.9. Subroutine UTimeInc (Overriding the Program-Determined Time Step)

This subroutine allows you to create a user-defined time step to override the one determined by the program. Activate the subroutine via the **USRCAL**,UTIMEINC command.

```
*deck,UTimeInc
                                       USERDISTRIB
      subroutine UTimeInc (deltmin,deltmax,delt)
                        User routine to override the program determined time step
c primary function:
С
                        Needs to have USRCAL, UTIMEINC to be accessed.
С
                        Called after the program determined the next time step
                          increment (AUTOTS,ON only)
C
           *** Copyright ANSYS. All Rights Reserved.
С
          *** ansys, inc.
C
С
  input arguments:
                (int,dp,in) - minimum time step size (user input)
(int,dp,in) - maximum time step size (user input)
      deltmin (int,dp,in)
C
С
               (int,dp,inout) - on input, the value determined by the program
C
c output arguments:
               (int,dp,inout) - on output, the value you have determined
```

2.3.10. Subroutine UCnvrg (Overriding the Program-Determined Convergence)

This subroutine allows you to create user-defined convergence checking and to override the convergence determined by the program. Activate the subroutine via the **USRCAL**, UCNVRG command.

```
*deck,UCnvrg USERDISTRIB
subroutine UCnvrg (ConvergenceType,ConvergenceFlag)

c primary function: User routine to perform custom convergence checking and
c override the program-determined convergence
c Needs to have USRCAL,UCNVRG to be accessed.
c Called after the program convergence checks.
```

```
*** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
C
c input arguments:
     ConvergenceType (int,sc,in)
                                      - type of convergence to be checked
C
C
                                          1, nonlinear element (called after
С
                                             element matrix formation)
                                          2, force convergence (called after
C
С
                                             element matrix formation)
                                          3, displacement convergence (called after
С
C
                                             equation solution)
С
      ConvergenceFlag (int,sc,inout) - on input, the value the program determined
                                        for this Type
C
С
                                          0, not converged
С
                                          1, converged
С
  output arguments:
     ConvergenceFlag (int,sc,inout) - on output, the value the you have determined
C
                                        for this Type
C
                                          0, not converged
C
                                          1, converged
С
      Note: For overall convergence, all 3 Types must be converged. Not all
С
            Types are evaluated (dependent on CNVTOL input and program defaults)
```

2.4. Subroutines for Customizing Material Behavior

This section describes the following subroutines that you can use to modify or monitor material behavior:

- 2.4.1. Subroutine UserMat (Creating Your Own Material Model)
- 2.4.2. Subroutine UserMatTh (Creating Your Own Thermal Material Model)
- 2.4.3. Subroutine UserHyper (Writing Your Own Isotropic Hyperelasticity Laws)
- 2.4.4. Subroutine UserHyperAniso (Writing Your Own Anisotropic Hyperelasticity Laws)
- 2.4.5. Subroutine UserCreep (Defining Creep Material Behavior)
- 2.4.6. Subroutine user_tbelastic (Defining Material Linear Elastic Properties)
- 2.4.7. Subroutine userfc (Defining Your Own Failure Criteria)
- 2.4.8. Subroutine userCZM (Defining Your Own Cohesive Zone Material)
- 2.4.9. Subroutine userswstrain (Defining Your Own Swelling Laws)
- 2.4.10. Subroutine userck (Checking User-Defined Material Data)
- 2.4.11. Supporting Function egen
- 2.4.12. Subroutine userfld (Update User-Defined Field Variables)
- 2.4.13. Subroutine userthstrain (Defining Your Own Thermal Strain)

Using the "_MATL" String

If you write a material-behavior subroutine using the **MPDATA**, **MPDELE**, **TB**, or **TBDELE** command, be aware that when the string "_MATL" appears in the MAT field of the command, the command interprets the string to mean the currently active material (as defined via the **MAT**, MAT command).

The "_MATL" string is used in conjunction with the library (LIB) option of the MPREAD and MPWRITE commands. When you issue MPWRITE with the LIB option, the command inserts "_MATL" in lieu of the specified material number as these commands are written to the material library file. When the program reads a material library file written in this format, it interprets "_MATL" to mean the currently active material. Do not use the "_MATL" string outside the scope of the MPREAD command.

2.4.1. Subroutine UserMat (Creating Your Own Material Model)

The UserMat subroutine allows you to write your own material constitutive equations within a general material framework using current-technology elements.

UserMat is a tool for advanced users. Expertise in material constitutive modeling and software programming is necessary. Developing a custom material constitutive model requires validation and testing. ANSYS, Inc. strongly recommends testing both single elements and multiple elements with various loading conditions to ensure correct results. UserMat supports shared memory and distributed parallel processing; however, you are responsible for ensuring that your code can use parallel processing.

The following UserMat topics are available:

- 2.4.1.1. UserMat Element Support
- 2.4.1.2. UserMat Overview
- 2.4.1.3. Stress, Strain, and Material Jacobian Matrix
- 2.4.1.4. The UserMat API
- 2.4.1.5. UserMat Variables
- 2.4.1.6. Table (TB) Commands for UserMat
- 2.4.1.7. Material Constitutive Integration with UserMat
- 2.4.1.8. UserMat Restrictions
- 2.4.1.9. Accessing Material and Element Data for UserMat
- 2.4.1.10. Utility Functions for UserMat

For a UserMat subroutine example, see Appendix C: User Material (UserMat) Subroutine Example (p. 375). For an example of UserMat in a coupled-field analysis, see Appendix D: Structural-Thermal User Material (UserMat, UserMatTh) Example (p. 385).

2.4.1.1. UserMat Element Support

Element support for user-defined material models (**TB**,USER) is available in Material Model Element Support in the *Material Reference*.

2.4.1.2. UserMat Overview

The UserMat subroutine defines the material stress-strain relationship of a material and applies to time-domain and full-harmonic analysis types. The subroutine supports current-technology elements only and does not apply to legacy elements.

The subroutine is called at every material integration point of the elements during the solution phase. The program passes in stresses, strains, and state variable values at the beginning of the

increment (and for time-domain analyses, the strain increment at the current increment). The UserMat subroutine then updates the stresses and state variables to current values.

2.4.1.2.1. Time Domain

The two most important UserMat-calculated quantities are stress and the consistent tangent stiffness:

- The measure of the stress is Cauchy stress (true stress), σ .
- The consistent tangent stiffness is defined as $\partial \Delta \sigma / \partial \Delta \varepsilon$, where $\Delta \sigma$ is the stress increment and $\Delta \varepsilon$ is the strain increment. The measure of strain is logarithmic strain (true strain), ε .

Generally, the material model can be formulated in either incremental (rate) or total forms.

- The incremental form is suitable for material models such as plasticity. The program uses a corotated framework for such material formulation. All variables are defined and updated in the co-rotated material coordinate system. The implementation of a constitutive model looks exactly the same for both small and large deformation (**NLGEOM**,ON).
- The total form is commonly used with hyperelastic material, and the material response is characterized by a potential relating to the invariants of the deformation gradient. The discrete equations are formulated with respect to the reference configuration. The deformation gradient is defined in the global coordinate system (with the exception of shell elements, where the deformation gradient is expressed on the co-rotated element coordinate system).

For most of the hyper-elastic models, material behavior is considered to be incompressible or nearly incompressible, resulting in a singular element matrix and possibly leading to element volumetric locking and solution failure. It is therefore necessary to take measures to avoid the singularity. The penalty method of carefully selecting the penalty parameter (which is also the compressibility parameter) is the simplest approach. Elements with mixed u-P formulation are the best way to address the singularity caused by material incompressibility. To use this option, set TBOPT = MXUP and element KEYOPT(6) = 1.

For a user-defined material with purely incompressible behavior, a mixed formulation in which the pressure is a degree of freedom is necessary. When a nearly incompressible hyperelastic material is formulated in UserMat, additional information (derivatives of volumetric potential with regard to determinant of deformation gradient J) is required. For more information, see UserMat Variables (p. 204).

Input values and the number of state variables (if used) for UserMat are specified via the **TB** command. For more information, see Table (**TB**) Commands for UserMat (p. 208).

2.4.1.2.2. Harmonic

For full-harmonic analyses, the incoming strain argument is a two-dimensional vector containing the real (material stiffness) and imaginary (damping) parts of the harmonic strain. The subroutine must calculate and return the real and imaginary parts of the stress and material Jacobian matrix for the current frequency.

The complex stress returned by the subroutine has no effect on the harmonic solution and is used for postprocessing only.

2.4.1.3. Stress, Strain, and Material Jacobian Matrix

For nonlinear geometry analysis, the stress measure (σ) used by the subroutine is the Cauchy stress (true stress), and the strain measure (ε) is the logarithmic strain (true strain). The strains and incremental strains passed into UserMat are the total mechanical strains from which the thermal strains (if they exist) are subtracted.

UserMat must also provide the material Jacobian matrix defined as $\partial \Delta \sigma_{ij} / \partial \Delta \varepsilon_{ij}$. $\Delta \sigma_{ij}$ is the stress increment, and $\Delta \varepsilon_{ii}$ is the strain increment.

UserMat is based on the current configuration for nonlinear geometry analysis (**NLGEOM**,ON). The program uses a co-rotational approach to account for rigid body rotation. Because the program already accounts for the strains passed into UserMat for the rigid body rotation, there is no need to apply additional rotation within UserMat.

Stress, strain, and the material Jacobian tensors are stored in a vector or matrix format.

The order of components for all tensors is as follows:

3-D stress state

11, 22, 33, 12, 23, 13

2-D plane strain and axisymmetric stress states

11, 22, 33, 12

2-D plane stress states

11, 22, 12

Beam element stress states

11, 13, 12

Link element stress state

11

The order of components for the material Jacobian matrix is as follows:

3-D stress state

1111	1122	1133	1112	1123	1113
2211	2222	2233	2212	2223	2213
3311	3322	3333	3312	3323	3313
1211	1222	1233	1212	1223	1213
2311	2322	2333	2312	2323	2313
1311	1322	1333	1312	1323	1313

2-D plane strain and axisymmetric stress states

1111	1122	1133	1112
2211	2222	2233	2212
3311	3322	3333	3312
1211	1222	1233	1212

2-D plane stress states

1111	1122	1112
2211	2222	2212
1211	1222	1212

Beam element stress states

1111	1113	1112
1311	1313	1312
1211	1213	1212

Link element stress state

1111

2.4.1.4. The UserMat API

Following is the interface for the UserMat subroutine:

```
*deck,usermat
                                    USERDISTRIB
     subroutine usermat(
                         matId, elemId,kDomIntPt, kLayer, kSectPt,
     &
                         ldstep, isubst, keycut,
     &
                         nDirect, nShear, ncomp, nStatev, nProp,
     &
                         Time, dTime, Temp, dTemp,
                          stress, ustatev, dsdePl, sedEl, sedPl, epseq,
                         Strain, dStrain, epsPl, prop, coords,
                          var0, defGrad_t, defGrad,
                          tsstif, epsZZ,
                         cutFactor, pVolDer, hrmflg, var3, var4,
                         var5, var6, var7)
      *** primary function ***
С
С
            user defined material constitutive model
C
      Attention:
С
            User must define material constitutive law properly
C
            according to the stress state such as 3D, plane strain
С
            and axisymmetry, plane stress and 3D/1D beam.
C
            A 3D material constitutive model can be used for
С
С
            plane strain and axisymmetry cases.
С
С
            When using shell elements, a plane stress algorithm
С
            must be used.
C
                                               gal July, 1999
```

```
C
С
       The following demonstrates a USERMAT subroutine for
       a plasticity model, which is the same as TB, BISO,
C
        for different stress states.
С
С
       See "ANSYS user material subroutine USERMAT" for detailed
       description of how to write a USERMAT routine.
C
С
       This routine calls four routines,
С
       usermat3d.F, usermatps.F usermatbm.F and usermat1d.F, w.r.t.
C
C
        the corresponding stress states.
       Each routine can be also a usermat routine for the specific
C
С
c Copyright ANSYS. All Rights Reserved.
С
C
      input arguments
С
      ==========
C
      matId
                 (int,sc,i)
                                         material #
      elemId
                (int,sc,i)
                                         element #
С
      kDomIntPt (int,sc,i)
                                         "k"th domain integration point
C
                (int,sc,i)
                                         "k"th layer
      kLayer
C
С
      kSectPt
                 (int,sc,i)
                                          "k"th Section point
      ldstep
                 (int,sc,i)
                                         load step number
С
                                         substep number
      isubst
                (int,sc,i)
C
      nDirect
                                         # of direct components
С
                 (int,sc,in)
      nShear
С
                 (int,sc,in)
                                         # of shear components
C
      ncomp
                 (int,sc,in)
                                         nDirect + nShear
                                         Number of state variables
С
      nstatev
                 (int,sc,i)
                (int,sc,i)
                                         Number of material constants
С
      nProp
С
С
      Temp
                 (dp,sc,in)
                                         temperature at beginning of
C
                                         time increment
                 (dp,sc,in)
      dTemp
                                         temperature increment
С
                                         time at beginning of increment (t)
С
      Time
                 (dp,sc,in)
      dTime
                (dp,sc,in)
                                         current time increment (dt)
C
С
С
      Strain (dp,ar(ncomp),i)
                                         Strain at beginning of time increment
      dStrain (dp,ar(ncomp),i)
                                         Strain increment
С
С
               (dp,ar(nprop),i)
                                         Material constants defined by TB, USER
      coords
                                         current coordinates
С
               (dp,ar(3),i)
С
      defGrad_t(dp,ar(3,3),i)
                                         Deformation gradient at time t
      defGrad (dp,ar(3,3),i)
                                         Deformation gradient at time t+dt
С
С
      hrmflg
               (dp,sc,io)
                                         flag to indicate harmonic analysis
С
С
      input output arguments
C
С
      stress (dp,ar(ncomp),io)
                                         stress
С
      ustatev (dp,ar(nstatev),io)
                                         user state variables
      sedEl
               (dp,sc,io)
                                         elastic work
C
      sedPl
               (dp,sc,io)
                                         plastic work
С
С
      epseq
               (dp,sc,io)
                                         equivalent plastic strain
С
      epsPl
               (dp,ar(ncomp),io)
                                         plastic strain
                                         not used, they are reserved arguments
С
      var?
               (dp,sc,io)
                                         for further development
C
С
С
      output arguments
C
      ______
                                         loading bisect/cut control
С
      keycut (int,sc,o)
С
                                         0 - no bisect/cut
                                         1 - bisect/cut
C
С
                                         (factor will be determined by solution control)
      dsdePl
                                         material jacobian matrix
С
                (dp,ar(ncomp,ncomp),o)
      pVolDer
                                         derivatives of volumetric potential wrt to J
C
               (dp,ar(3),o)
                                         pVolDer(1) = dU/dJ
C
                                         pVolDer(2) = d^2U/dJ^2
C
                                         pVolDer(3) = d^3U/dJ^3
С
С
      tsstif
                (dp,ar(2),o)
                                         transverse shear stiffness
                                         tsstif(1) - Gxz
С
                                         tsstif(2) - Gyz
С
                                         tsstif(1) is also used to calculate hourglass
```

```
С
                                      stiffness, this value must be defined when low
С
                                     order element, such as 181, 182, 185 with uniform
С
                                     integration is used.
С
      epsZZ
                                     strain epsZZ for plane stress,
              (dp,sc,o)
С
                                     define it when accounting for thickness change
                                     in shell and plane stress states
C
      cutFactor(dp,sc,o)
                                      time step size cut-back factor
С
                                     define it if a smaller step size is wished
С
                                     recommended value is 0~1
C
C'
С
С
      ncomp
             6
                 for 3D (nshear=3)
                 for plane strain or axisymmetric (nShear = 1)
С
      ncomp
             4
      ncomp
С
                 for plane stress (nShear = 1)
С
      ncomp
            3 for 3d beam (nShear = 2)
                for 1D (nShear = 0)
            1
С
      ncomp
С
С
      stresses and strains, plastic strain vectors
         11, 22, 33, 12, 23, 13 for 3D
С
         11, 22, 33, 12
                                 for plane strain or axisymmetry
C
         11, 22, 12
                                 for plane stress
С
С
         11, 13, 12
                                 for 3d beam
С
          11
                                 for 1D
C
      material jacobian matrix
С
С
        3D
                                        1112
C
          dsdePl
                     1111
                           1122 1133
                                              1123
                                                     1113
                      2211
                             2222
                                   2233
                                                      2213
С
          dsdePl
                                         2212
                                               2223
                           3322
                                  3333
                      3311
                                                      3313
С
          dsdePl
                                         3312
                                               3323
          dsdePl
                     1211 1222 1233 1212
                                              1223
                                                     1213
С
          dsdePl
                   2311 2322 2333 2312
                                               2323
                                                      2313
С
С
          dsdePl
                   | 1311 1322 1333 1312
                                              1323
                                                     1313
        plane strain or axisymmetric (11, 22, 33, 12)
С
С
          dsdePl
                     1111 1122
                                   1133
                                        1112
          dsdePl
                      2211 2222
                                  2233
                                        2212
C
С
          dsdePl
                     3311 3322 3333 3312
С
          dsdePl
                    1211 1222 1233 1212
        plane stress (11, 22, 12)
С
С
          dsdePl
                     1111
                            1122
                                   1112
                      2211
                            2222
                                   2212
С
          dsdePl
          dsdePl
                    1211 1222
                                  1212
С
        3d beam (11, 13, 12)
С
С
          dsdePl | 1111
                            1113
                                  1112
С
          dsdePl
                      1311
                            1313
                                   1312
С
          dsdePl
                    1211
                            1213
                                   1212
        1d
С
С
          dsdePl
                   | 1111 |
```

The ncomp value used in get_ElmData (p. 210) and put_ElmData (p. 383) may differ from the ncomp value passed into usermat.F. Use get_ElmData with the 'NCOMP' option to determine the correct array sizes.

2.4.1.5. UserMat Variables

The UserMat subroutine uses the following Input (p. 204), Input/Output (p. 206), and Output (p. 207) variables. Do not change them in the subroutine code.

	UserMat Input Arguments
matld	Integer variable containing the material ID number.
elemId	Integer variable containing the element number.

kDomIntPt	Integer variable containing the material integration point number.
kLayer	Integer variable containing the layer number.
kSectPt	Integer variable containing section point number.
ldstep	Integer variable containing load step number.
isubst	Integer variable containing substep number.
nDirect	Number of direct components of the stress or strain vector at material point.
nShear	Number of shear components of the stress or strain vector at material point (engineering components).
ncomp	Total number of the stress or strain components at material point (nDirect + nShear).
nstatev	Number of state variables, specified by the $NPTS$ value in the TB ,STATE command.
nProp	Number of material constants, specified by the $NPTS$ value in the TB ,USER command
Temp	Double-precision variable containing the temperature at the beginning of time increment.
dTemp	Double-precision variable containing the current temperature increment.
Time	For time-domain analyses, a double-precision variable containing the total time at the beginning of the time increment. For full-harmonic analyses, the current value of frequency.
dTime	For time-domain analyses, a double-precision variable containing the current time increment. For full-harmonic analyses, the frequency increment.
Strain	Double-precision array containing the total strains at the beginning of the time increment. Array size is Strain(ncomp) in time-domain analyses and Strain(ncomp,2) in harmonic analyses, where the real and imaginary strain components are in the first and second column, respectively.
	Thermal strains (defined via MP ,ALPHA and temperature load), if any, are subtracted from the total strains; therefore, the strains passed to UserMat are the mechanical strains only.
	For large-deformation problems, (NLGEOM ,ON), the strain components are updated to account for rigid body rotation before they are passed to UserMat and are approximately the logarithmic strains.
	When the mixed u-P formulation option (TBOPT = MXUP on the TB ,USER command) is used for hyperelastic material, the strain array is the logarithmic strains at the current time. However, the strain array can be redefined within the UserMat subroutine. For nearly incompressible hyperelastic material, a mixed u-J formulation is used. The calculated J is passed from strain array as strain(ncomp+1).
dStrain	Double-precision array containing current strain increments. Array size is ncomp. As with the Strain array, this value contains the mechanical strain increments only. Thermal strain increments (if any) are subtracted from the total strains increments.
	When the mixed u-P formulation option ($TBOPT = MXUP$ on the TB ,USER command) is used for hyperelastic material, the dStrain array is zero.

prop	Double-precision array containing the material constants defined via TB ,USER and TBDATA commands. Array size is nProp. Array prop contains the material constants at current temperature point.
coords	Double-precision array containing the current coordinates of the material integration points. Array size is 3.
defGrad_t	Double-precision matrix containing deformation gradient at the beginning of the time increment. The matrix size is 3 x 3. The matrix components $DefGrad_{(i,j)}$ are equivalent to deformation gradient F_{ij} at the beginning of the time increment and are only available for continuum and shell elements with nonlinear deformation (NLGEOM ,ON).
defGrad	Double-precision matrix containing current deformation gradient. The matrix size is 3 x 3. The matrix components $\operatorname{DefGrad}_{(i,j)}$ are equivalent to deformation gradient F_{ij} at the current time and are only available for continuum and shell elements with nonlinear deformation (NLGEOM ,ON).

UserMat Input/Output Arguments		
stress	Double-precision array containing the stresses.	
	For time-domain analyses, its size is defined by the ncomp input value. The stress measure is the "true" stress. It is passed as the values of stresses at the beginning of the time increment and must be updated to the values of stress at the end of the time increment.	
	For harmonic analyses, its size is stress(ncomp,2), where the first column is the real stress and the second column is the imaginary stress.	
	For finite-deformation problems, the stresses are rotated to account for rigid body motion before they are passed in, and therefore only the co-rotational portion of stress integration is required in UserMat.	
	When the mixed u-P formulation option ($TBOPT = MXUP$ on the TB ,USER command) is used for hyperelastic material, the stress is updated for deviatoric part of stress only. The calculated P is passed into usermat as stress(ncomp+1).	
statev	Double-precision array containing the state variables. Its size is defined via the TB ,STATE command. It is passed as the values of state variables at the beginning of the time increment and must be updated to the values of the state variables at the end of the time increment.	
epseq	Equivalent plastic strain.	
epspl	Double-precision array containing the plastic strains. The strain measure is the "true" strain. Its size is defined by the ncomp input value. It is passed as the values of the plastic strains at the beginning of the time increment and must be updated to the values of the plastic strains at the end of the time increment.	
	For finite-deformation problems, the plastic strains have been rotated to account for rigid body motion before they are passed in.	
sedEl	Elastic work. It is used for output purposes only and does not affect the solution.	
sedPl	Plastic work. It is used for output purposes only and does not affect the solution.	

UserMat Output Arguments		
	These values must be updated in the subroutine code.	
keycut	Integer variable as key for loading bisection/cut control:	
	0 - No bisect/cut (default)	
	1 - Bisect/cut	
	Set keycut = 1 when UserMat experiences convergence difficulty when solving constitutive equation integration. The bisect/cut factor is determined by the solution control. Set cutFactor to control the time stepping size.	
	Not used in harmonic analyses.	
epsZZ	Strain component at an out-of-plane direction for the plane stress state. This value is required when the thickness change is taken into account in plane stress or shell elements.	
tsstif(2)	Transverse shear stiffness:	
	Tsstif(1) - GXZ	
	Tsstif(2) - GYZ	
dsdePl	Double-precision array containing the material Jacobian matrix	
	$\partial \Delta \sigma_{ij} / \partial \Delta \varepsilon_{ij}$. Here, the values represent the stress/strain increments, respectively. The dsdePl(i,j) value denotes the change in the i-th stress component caused by a change of the j-th strain component.	
	By default, the program assumes that the element stiffness matrix is symmetric; therefore, you must provide a <i>symmetric</i> material Jacobian matrix <i>even if it is unsymmetric</i> . If your material requires an unsymmetric material Jacobian matrix, issue the NROPT , UNSYM command to define the unsymmetric stiffness matrix.	
	When the mixed u-P formulation option ($TBOPT = MXUP$ on the TB ,USER command) is used for hyperelastic material, only the deviatoric material consistent tangent matrix is needed.	
	For harmonic analyses, the real components are returned in dsdePl(1:ncomp,1:ncomp,1), and the imaginary components are returned in dsdePl(1:ncomp,1:ncomp,2).	
pVolDer(:)	Derivatives of volumetric potential with regard to the determinant of deformation gradient:	
	$pVolDer(1) = \frac{\partial U}{\partial J}$	
	$pVolDer(2) = \frac{\partial^2 U}{\partial J^2}$	
	$pVolDer(3) = \frac{\partial^3 U}{\partial J^3}$	

	This argument is needed only when a hyperelastic material is defined and mixed u-P formulation is used.
	For nearly incompressible hyperelastic material, a mixed u-J formulation is used. The derivatives of volume potential is to Jc, where Jc is the calculated J and is passed from strain array as strain(ncomp+1).
	For purely incompressible hyperelastic material, set all three derivatives pVolDer(1:3) to zero.
cutFactor	Time-step size control factor.
	Not used in harmonic analyses.

2.4.1.6. Table (TB) Commands for UserMat

When creating your own material model, first define the material by specifying input values for the UserMat subroutine (**TB**,USER). It is also necessary to specify the number of state variables used, if applicable (**TB**,STATE).

Following is more information about defining your material and specifying the number of state variables used. For detailed information about the **TB** command and arguments, see the *Command Reference*.

TB, USER Command

Issue the **TB** command using the following syntax:

```
TB, USER, MAT, NTEMPS, NPTS, TBOPT
```

where

MAT = User material ID number

NTEMPS = Number of temperature points.

NPTS = Number of material constants at a given temperature point.

TBOPT = NONLINEAR (default), LINEAR, or MXUP

The material properties at an intermediate temperature point are interpolated and passed to the UserMat subroutine.

Define temperatures and material constants via **TBTEMP** and **TBDATA** commands, respectively.

Example 2.1: Defining the Material for UserMat

```
tb,user,1,2,4
! Define material 1 as a user
! material with two temperatures
! and four data points at each
! temperature point.

tbtemp,1.0
! first temp.

tbdata,1,19e5, 0.3, 1e3,100, ! Four mat. constants for one temp.

tbtemp,2.0
! Second temp.

tbdata,1,21e5, 0.3, 2e3,100, ! Four mat. constants for two temps.
```

TB,STATE Command

If you intend to use state variables with the UserMat subroutine, it is necessary to first specify the number of state variables. Issue the **TB** command using the following syntax:

```
TB,STATE,MAT, ,NPTS

where

MAT = User material ID number

NPTS = Number of state variables that you intend to use.
```

The command defines only the *number* of state variables and must always be associated with a user material ID. No temperatures or data are associated with the command.

By default, the program initializes state variables to zero at the beginning of an analysis. Use the **TBDATA** command to initialize your own values for state variables.

Example 2.2: Defining the Number of State Variables for UserMat

```
tb,state,1,,8 ! Define material 1 with eight state variables tbdata,1,c1,c2,c3,c4,c5,c6,c7,c8 ! Initialize the eight state variables.
```

2.4.1.7. Material Constitutive Integration with UserMat

The UserMat subroutine supports current-technology elements with all key options. However, a different material constitutive integration is necessary for the various stress states, such as general 3-D, plane stress, and beam (with or without shear-stress components).

To ensure overall numerical stability, verify that the integration scheme implemented in the subroutine is stable. The program always uses the full Newton-Raphson scheme for the global time-domain solution to achieve a better convergence rate. The material Jacobian matrix (p. 201) (dsde-Pl(i,j) (p. 207)) must be consistent with the material constitutive integration scheme for a better convergence rate of the overall Newton-Raphson scheme.

2.4.1.8. UserMat Restrictions

The following restrictions apply to the UserMat subroutine:

- The subroutine supports current-technology elements only and does not apply to legacy elements.
 - For more information, see Older vs. Current Element Technologies in the *Element Reference*.
- The state variables (defined via the **TB**,STATE command) are supported only by full graphics in the POST1 postprocessor.
 - Because POST1 does not switch to full graphics automatically, you must issue a /GRAPHICS, FULL command to do so.
- The UserMat interface for elements with mixed u-P formulation is supported only for current-technology continuum elements (such as PLANE182, PLANE183, SOLID185, SOLID186, SOLID187, SOLSH190, CPT212, CPT213, CPT215, CPT216, CPT217, SOLID226, and SOLID227), excluding plane

stress state. You must set KEYOPT(6) = 1 with elements PLANE182, PLANE183, SOLID185, SOLID186, SOLID187, SOLSH190, CPT212, CPT213, CPT215, CPT216, and CPT217 and KEYOPT(11) = 1 with SOLID226 and SOLID227 . For elements with a plane stress assumption, a mixed u-P formulation is not needed.

 The deformation gradient defGrad(3,3) and defGrad_t(3,3) are available only for continuum and shell elements with nonlinear deformation (NLGEOM,ON).

2.4.1.9. Accessing Material and Element Data for UserMat

Following is the interface for accessing the material and element data:

```
*deck, get ElmData
     subroutine get_ElmData (kchar, elemId, kMatRecPt, ncomp, vect)
C
С
     *** primary function
С
          retrieve material record data
С
С
          including items:
             stress vector
C
C
             elastic strain vector
C
             plastic strain vector
C
             creep strain vector
С
              thermal strain vector
C
             state variables
С
С
                                 ---- Guoyu Lin 4/25/2001 ----
C
c *** Notice - This file contains ANSYS Confidential information ***
С
С
    input arguments
С
    ==========
С
    kchar (ch,sc,in)
                               string characters indicating
                               inquired quantities
С
С
    kMatRecPt
               (in,sc,in)
                              integration point where data
С
                              to be inquired
             (in,sc,in) element number (in,sc,in) number of compo
    elemId
C
С
    ncomp
                               number of components to be inquired
С
                               Use the 'NCOMP' query to get the right
                               size for tensor quantities.
C
С
    numWordsPt (in,ar(*),in)
                              number of mat. Record
С
С
    input output arguments
                              input desc / output desc
     _____
                              ========
                                            _____
С
С
С
    output arguments
C
           (dp,ar(*),ou) inquired data
C
    vect.
С
С
    local variables
С
     ==========
c --- parameters
```

The ncomp value used in get_ElmData (p. 210) and put_ElmData (p. 383) may differ from the ncomp value passed into usermat.F (p. 202). Use get_ElmData with the 'NCOMP' option to determine the correct array sizes.

For a descriptions of the input arguments and valid argument variables, see Accessing Solution and Material Data (p. 383).

2.4.1.10. Utility Functions for UserMat

The following functions are available for use with UserMat. ANSYS, Inc. provides them for your convenience.

	Utility Functions for UserMat		
vzero(a ,n)	Initializes array \mathbf{a} to zero. The value n is the array dimension.		
vmult(a , b ,n,c)	Multiplies vector \mathbf{a} by constant c and outputs ($\mathbf{b} = \mathbf{a} * c$) as vector \mathbf{b} . The value n is the dimension for both arrays.		
vmult1(a ,n,c)	Multiplies vector \mathbf{a} by constant \mathbf{c} and outputs the result as itself (that is, $\mathbf{a} = \mathbf{a} * \mathbf{c}$). The value \mathbf{n} represents the array dimension.		
maxb(a, b , c, na, nb, nc,	Multiplies two double-precision matrices and outputs the result as c (that is, $c = a * b$). The value na is number of rows in matrix a , nb the number of rows in matrix b , and nc the number of rows in matrix c .		
n1, n2, n3)	The $n1$ value is the number of rows in matrix c to fill, and $n2$ the number of columns in matrix c to fill. The value $n3$ is the number of columns in matrix c and the number of rows in matrix c to work with. (The number of columns in c and rows in c is the same in order to generate the inner product correctly.)		

2.4.2. Subroutine UserMatTh (Creating Your Own Thermal Material Model)

UserMatTh is a tool for advanced users. Expertise in thermal material modeling and software programming is necessary. Developing a custom thermal material model requires validation and testing. ANSYS, Inc. strongly recommends testing both single elements and multiple elements with various loading conditions to ensure correct results. UserMatTh supports shared memory and distributed parallel processing; however, you are responsible for ensuring that your code can use parallel processing.

The following UserMatTh topics are available:

- 2.4.2.1. UserMatTh Element Support
- 2.4.2.2.The UserMatTh API
- 2.4.2.3. UserMatTh Variables
- 2.4.2.4. Table (TB) Commands for UserMatTh
- 2.4.2.5. UserMatTh Restrictions
- 2.4.2.6. Utility Functions for UserMatTh

2.4.2.1. UserMatTh Element Support

Element support for user-defined thermal material models (**TB**,USER) is available in Material Model Element Support in the *Material Reference*.

UserMatTh is called at every material integration point of the elements during the solution phase.

Input values and the number of state variables (if used) for the subroutine are specified via **TB**. For more information, see Table (**TB**) Commands for UserMatTh (p. 215).

2.4.2.2. The UserMatTh API

Following is the interface for the UserMatTh subroutine:

```
*deck,usermatth
                                 USERDISTRIB
    subroutine usermatth(matId, elemId, kDomIntPt, kLayer, kSectPt,
                        ldstep, isubst, keycut, ncomp, nStatev,nProp,
                        Time, dTime, Temp, dTemp, tgrad,
    S.
                        ustatev, prop, coords,
                        dudt, dudg, flux, dfdt, dfdg,
                        cutFactor, hgen, dens, var1, var2, var3,
                        var4, var5, var6)
*** primary function ***
С
С
          user defined thermal material constitutive model
С
С
    Attention:
C
          User must define material constitutive behavior properly
С
С
С
С
       The following demonstrates a USERMATTH subroutine for
С
       a regular conductive heat transfer.
       See "ANSYS user material subroutine USERMATTH" for detailed
С
С
       description of how to write a USERMATTH routine.
C
С
c Copyright ANSYS. All Rights Reserved.
C
С
     input arguments
     ==========
C
      matId
С
     elemId (int,sc,i)
                                     element #
С
С
     kDomIntPt (int,sc,i)
                                     "k"th domain integration point
С
     kLayer (int,sc,i)
                                     "k"th layer
     kSectPt (int,sc,i)
                                     "k"th Section point
С
     ldstep
С
              (int,sc,i)
                                     load step number
С
     isubst
              (int,sc,i)
                                     substep number
С
     ncomp
              (int,sc,i)
                                     # of components
                                     Number of state variables
С
     nStatev (int,sc,i)
С
     nProp
              (int,sc,i)
                                     Number of material constants
C
      Temp
               (dp,sc,in)
                                     temperature at current time
С
С
      dTemp
               (dp,sc,in)
                                     temperature increment
С
      Time
               (dp,sc,in)
                                     time at beginning of increment (t)
С
      dTime
               (dp,sc,in)
                                     current time increment (dt)
С
С
      prop
               (dp,ar(nprop),i)
                                     Material constants defined by TB, USER
      coords
С
               (dp,ar(3),i)
                                     current coordinates
C
               (dp,ar(ncomp),i)
                                     Current values of the spatial gradients of temperature
С
С
С
     input output arguments
```

```
С
     -----
                                      user state variables
С
      ustatev (dp,ar(nstatev),io)
                (dp,sr,io)
                                      heat generation rate per unit mass
C
      hqen
С
      dens
               (dp,sr,io)
                                       density passed in as defined mp command
С
      var?
                (dp,sc,io)
                                       not used, they are reserved arguments
                                       for further development
С
С
С
     output arguments
C
     ==========
С
      keycut (int,sc,o)
                                       loading bisect/cut control
                                       0 - no bisect/cut
С
                                       1 - bisect/cut
С
С
                                       (factor will be determined by ANSYS solution control)
C
      cutFactor (dp,sc,o)
                                       time step size cut-back factor
С
                                       define it if a smaller step size is wished
С
                                       recommended value is 0~1
С
С
С
      dudt
                (dp,sr,o)
                                       unit mass internal energy variation with respect to
С
                                       temperature
С
      dudg
                (dp,ar(ncomp),o)
                                       unit mass internal thermal energy variation with
                                      respect to the spatial gradients of temperature
C
С
      flux
                (dp,ar(ncomp),o)
                                      heat flux vector
С
      dfdt
                (dp,ar(ncomp),o)
                                       heat flux vector variation with respect to temperature
      dfda
                (dp,ar(ncomp,ncomp),o)
                                      heat flux vector variation with respect to the spatial
C
                                       gradients of temperature
С
С
С
                for 3D
С
      ncomp
                for 2D
С
      ncomp
С
      ncomp 1 for 1D
С
С
```

2.4.2.3. UserMatTh Variables

The UserMatTh subroutine uses the following Input (p. 213), Input/Output (p. 214), and Output (p. 214) variables. Do not change the Input arguments in the subroutine code.

	UserMatTh Input Arguments
matld	Integer variable containing the material ID number.
elemId	Integer variable containing the element number.
kDomIntPt	Integer variable containing the material integration point number.
kLayer	Integer variable containing the layer number.
kSectPt	Integer variable containing section point number.
ldstep	Integer variable containing load step number.
isubst	Integer variable containing substep number.
ncomp	Integer variable containing spatial dimension number: 1 for 1-D, 2 for 2-D, or 3 for 3-D.
nStatev	Number of state variables, specified via the \mathtt{NPTS} value in the \mathtt{TB} ,STATE command.
nProp	Number of material constants, specified via the NPTS value in the \textbf{TB} ,USER command
Temp	Double-precision variable containing the current temperature.
dTemp	Double-precision variable containing the current temperature increment.

Time	Double-precision variable containing the total time at the beginning of the time increment.	
dTime	Double-precision variable containing the current time increment.	
prop	Double-precision array containing the material constants defined via TB ,USER and TBDATA commands. Array size is nProp. Array prop contains the material constants at current temperature point.	
coords	Double-precision array containing the current coordinates of the material integration points. Array size is 3.	
tgrad	Double-precision array of size ncomp containing current temperature gradient.	

UserMatTh Input/Output Arguments		
hgen	Double-precision variable with heat-generation information. It is passed in as the value that is predefined with the model. The value can be updated with user-defined heat generation. [1] (p. 214)	
ustatev	Double-precision array containing the state variables. Its size is defined via the TB ,STATE command. It is passed as the values of state variables at the beginning of the time increment and must be updated to the values of the state variables at the end of the time increment. [1] (p. 214)	
dens	Double precision variable with density information. It is passed in as the value that is predefined with the model. The value can be updated with user-defined density. [1] (p. 214)	

- 1. In a coupled structural-thermal analysis using SOLID226 and SOLID227:
 - the heat generation rate (hgen) is interpreted as heat flow rate per unit volume,
 - the state variables (ustatev) cannot be updated with user-defined values in UserMatTh,
 - the density (dens) cannot be updated with user-defined value in UserMatTh.

UserMatTh Output Arguments			
These values must be updated in the subroutine code.			
keycut	Integer variable as key for loading bisection/cut control:		
	0 - No bisect/cut (default)		
	1 - Bisect/cut		
	Set keycut = 1 when UserMatTh experiences convergence difficulty when solving constitutive equation integration. The bisect/cut factor is determined by the solution control.		
cutFactor	Double-precision variable containing the user-defined time-step size-control factor.		
dudt	Double-precision variable containing the unit mass internal energy variation with respect to temperature.		

dudg	Double-precision array of size (ncomp,ncomp) containing the unit mass internal thermal energy variation with respect to the spatial gradients of temperature.	
flux	Double-precision array of size ncomp containing the heat flux vector.	
dfdt	Double-precision array of size ncomp containing the heat flux vector variation with respect to temperature.	
dfdg	Double-precision array of size (ncomp,ncomp) containing the heat flux vector variation with respect to the spatial gradients of temperature.	

2.4.2.4. Table (TB) Commands for UserMatTh

When creating your own material model, first define the material by specifying input values for the UserMatTh subroutine (**TB**,USER). It is also necessary to specify the number of state variables used, if applicable (**TB**,STATE).

In a coupled-field analysis with structural and thermal degrees of freedom, use **TB**,USER with *TBOPT* = THERM to define the thermal material independently of the structural material model. The THERM option is available with elements SOLID226 and SOLID227.

Following is more information about defining your material and specifying the number of state variables used. For detailed information about the **TB** command and arguments, see the *Command Reference*.

TB, USER Command

Issue the **TB** command using the following syntax:

```
TB, USER, MAT, NTEMPS, NPTS, TBOPT
```

where

MAT = User material ID number

NTEMPS = Number of temperature points.

NPTS = Number of material constants at a given temperature point.

TBOPT = NONLINEAR (default), LINEAR, or THERM

The material properties at an intermediate temperature point are interpolated and passed to the UserMatTh subroutine.

Define temperatures and material constants via **TBTEMP** and **TBDATA** commands, respectively.

Example 2.3: Defining the Material for UserMatTh

```
tb,user,1,2,4
! Define material 1 as a user
! material with two temperatures
! and four data points at each
! temperature point
tbtemp,1.0
! first temperature
tbdata,1,19e5,0.3,1e3,100,
! four material constants for
! one temperature
tbtemp,2.0
! Second temperature
```

```
tbdata,1,21e5,0.3,2e3,100, ! Four material constants for ! two temperatures
```

For an example of UserMatTh in a coupled-field analysis, see Appendix D: Structural-Thermal User Material (UserMat, UserMatTh) Example (p. 385).

TB,STATE Command

If you intend to use state variables with the UserMatTh subroutine, it is necessary to first specify the number of state variables. Issue the **TB** command using the following syntax:

TB,STATE,MAT, NPTS where MAT = User material ID number. NPTS = Number of state variables that you intend to use.

The command defines only the *number* of state variables and must always be associated with a user material ID. No temperatures or data are associated with the command.

By default, the program initializes state variables to zero at the beginning of an analysis. Use the **TBDATA** command to initialize your own values for state variables.

2.4.2.5. UserMatTh Restrictions

The following restrictions apply to the UserMatTh subroutine:

- The subroutine supports current-technology elements only and does not apply to legacy elements. For more information, see UserMatTh Element Support (p. 212).
- The state variables (defined via the **TB**,STATE command) are supported only by full graphics in the POST1 postprocessor.

Because POST1 does not switch to full graphics automatically, you must issue a /GRAPHICS, FULL command to do so.

• Element FLUID116 is supported only when KEYOPT(1) = 1. For this element, you can use User-MatTh to specify material properties KXX, C, and DENS only.

2.4.2.6. Utility Functions for UserMatTh

The following functions are available for use with UserMatTh. ANSYS, Inc. provides them for your convenience.

Utility Functions for UserMatTh		
1 . `	Initializes array a to zero.	
a ,n)	The value n is the array dimension.	

vmult(Multiplies vector \mathbf{a} by constant \mathbf{c} and outputs ($\mathbf{b} = \mathbf{a} * \mathbf{c}$) as vector \mathbf{b} .		
a , b ,n,c)	The value n is the dimension for both arrays.		
vmult1(a ,n,c)	Multiplies vector \mathbf{a} by constant \mathbf{c} and outputs the result as itself (that is, $\mathbf{a} = \mathbf{a} * \mathbf{c}$). The value \mathbf{n} represents the array dimension.		
maxb(a , b	Multiplies two double-precision matrices and outputs the result as c (that is, $c = a * b$).		
, c , na, nb, nc, n1,	The value na is number of rows in matrix a , nb the number of rows in matrix b , and nc the number of rows in matrix c . The $n1$ value is the number of rows in matrix c to fill, and $n2$ the number of columns in matrix c to fill.		
n2, n3)	The value $n3$ is the number of <i>columns</i> in matrix a and the number of <i>rows</i> in matrix b to work with. (The number of columns in a and rows in b is the same in order to generate the inner product correctly.)		

2.4.3. Subroutine UserHyper (Writing Your Own Isotropic Hyperelasticity Laws)

Use the subroutine UserHyper when you issue TB,HYPER with TBOPT = USER.

```
*deck, UserHyper
                                  USERDISTRIB
c Copyright ANSYS. All Rights Reserved.
     subroutine UserHyper(
    &
                          prophy, incomp, nprophy, invar,
    &
                          potential, pInvDer)
     *** Example of user hyperelastic routine
С
C
           This example uses Arruda hyperelasticity model
С
           which is the same ANSYS TB, HYPER, , , , BOYCE
С
C
     input arguments
    ==========
С
                  (dp,ar(*),i)
С
     prophy
                                  material property array
                 (int,sc,i)
С
      nprophy
                                  # of material constants
                dp,ar(3)
                                  invariants
      invar
С
С
С
     output arguments
C
     ==========
      incomp (log,sc,i)
                                  fully incompressible or compressible
С
     potential
                                   value of potential
                dp,sc
С
                   dp,ar(10)
                                   der of potential wrt i1,i2,j
     pInvDer
C
С
                                    1 - der of potential wrt il
                                    2 - der of potential wrt i2
C
                                    3 - der of potential wrt ilil
С
                                    4 - der of potential wrt ili2
С
                                    5 - der of potential wrt i2i2
C
                                    6 - der of potential wrt ilj
С
                                    7 - der of potential wrt i2j
С
                                    8 - der of potential wrt j
С
С
                                    9 - der of potential wrt jj
С
             ***********
```

```
c
c --- parameters
c
```

2.4.4. Subroutine UserHyperAniso (Writing Your Own Anisotropic Hypere-lasticity Laws)

The UserHyperAniso subroutine defines the potential derivatives for a strain-energy potential that is a function of isochoric strain invariants and anisotropic fiber invariants.

2.4.4.1. Input Parameters

Use the UserHyperAniso subroutine when you issue TB, AHYPER with TBOPT = USER.

Issue **TBDATA** to specify the invariant set type (SetType). The only valid set type is 101, which defines the invariant set as the isochoric strain invariants and isochoric fiber invariants defined below.

Example 2.4: User-Defined Anisotropic Hyperelastic Material Model

```
/prep7
TB,AHYPER,1,,,USER
TBDATA,1,101 ! define the invariant set type
```

You can define an optional set number (SetNumber) in a data table initiated via TBOPT = UNUM. This value is passed into the subroutine for use in distinguishing between user-defined material behaviors.

Example 2.5: User-Defined Anisotropic Hyperelastic Set Number

```
/prep7
TB,AHYPER,1,,,UNUM
TBDATA,1,1 ! define the invariant set number
```

Define the material parameters in a table initiated via TBOPT = AU01.

Example 2.6: User-Defined Anisotropic Hyperelastic Parameters

```
/prep7
TB, AHYPER, 1,,, AU01
 !a1, a2, a3
    tbdata,1,a1
 !b1, b2, b3
    tbdata,4,b1
 !c1, c2, c3,
    tbdata,7,c1, c2, c3,
 !d2, d3, d4, d5, d6
    tbdata, 12, 0, 0.0, 0.0, 0.0, 0.0
 !e1, e2, e3
    tbdata, 17, e1, e2, e3
 !f1, f2, f3, f4, f5, f6
    tbdata, 22, 0, 0.0, 0.0, 0.0, 0.0
 !g1, g2, g3, g4, g5, g6
    tbdata,27,0,0.0,0.0,0.0,0.0
    tbdata, 31, 1e-5
```

Specify the fiber directions in a data table with TBOPT = FB01. Three values define the direction of a fiber. The direction is relative to the element coordinate system and should have a magnitude equal to 1.0. The number of fibers (25 maximum) is determined from the number of defined values.

Example 2.7: User-Defined Anisotropic Hyperelastic Fiber Directions

```
/prep7
TB,AHYPER,1,,,FB01
TBDATA,1,1,0,0
TBDATA,4,1/sqrt(2),1/sqrt(2),0
```

2.4.4.2. Invariants and Potential Derivatives

Define the strain-energy potential as a function of the isochoric strain invariants and the anisotropic invariants that depend on the fiber directions and deformation.

$$W = W(\overline{I}_1, \overline{I}_2, J, A_1, \ldots, A_n)$$

where I_1, I_2, J are the isochoric strain invariants and $A_1, \blacksquare \blacksquare \blacksquare, A_n$ are the fiber directions. The strain-energy potential depends on isochoric fiber invariants that have the form:

$$\begin{split} & \bar{I}_4 = A_i \cdot \overline{C} A_i \\ & \bar{I}_5 = A_i \cdot \overline{C}^2 A_i \\ & \bar{I}_6 = A_i \cdot \overline{C} A_i, \quad i \neq j \end{split}$$

where I_4 is a first-order fiber invariant, I_5 is a second-order fiber invariant, and I_6 is a mixed-fiber variant.

Mixed second-order fiber invariants are not used in the strain-energy potential definition.

The isochoric strain invariants are numbered 1, 2, and 3 respectively in the Invar(*) array. The fiber-invariant numbering scheme is obtained from the following function:

```
ni = Flindx(SetType,InvType,Fib1,Fib2)
```

where:

```
ni = returned value of invariant number in Invar(*) array
SetType = set type passed into UserHyperAniso
InvType = invariant type: 1 = first-order, 2 = second-order
Fib1 = first fiber number, i
Fib2 = second fiber number, i or j
```

The UserHyperAniso subroutine should calculate and return the first and second derivatives of the user-defined strain-energy potential with respect to the invariants. For mixed u-P formulations that are not incompressible, the third-order invariant with respect to J is also required. (Third-order derivatives of the strain-energy potential with respect to other invariants are not required.)

The compressibility of the material is determined from the second derivative of the strain-energy potential with respect to J in the reference configuration during the first solution step. If this derivative is zero, the material is incompressible and the appropriate mixed u-P formulation is used. The material should not change between compressible and incompressible behavior during the simulation.

The strain-energy-potential derivatives are stored in the arrays pD1(*), pD2(*) and pD3(*) via the following subroutine:

```
put_PDer(SetType,nFib,Order,In1,In2,In3,pD1,pD2,pD3,val)
```

where:

```
SetType = set type passed into UserHyperAniso
nFib = number of fibers passed into UserHyperAniso
Order = derivative order to be stored (1, 2, or 3)
In1, In2, In3 = invariant numbers corresponding to the respective derivatives
pD1, pD2, pD3 = potential derivative arrays passed into UserHyperAniso
val = value of the derivative that is added to the value in the array
```

2.4.4.3. UserHyperAniso API

```
*deck, User Hyper Aniso
                                               USERDISTRIB
      subroutine UserHyperAniso(SetType, SetNumber, incomp, upkey, nprophy,
                             prophy,nFib,fibDir,ninv,Invar,potential,
     8
                              pD1,pD2,pD3)
С
      *** Example of user anisotropic hyperelastic routine
С
С
С
             This example reproduces the Polynomial hyperelasticity
            model which is the same as TB, AHYPER, , , , POLY
С
C
С
      input arguments
С
      _____
                                       Type of invariant set
                    (int,sc,i)
       SetType
C
       SetNumber (int,sc,i)
User input invariant set incomp (log,sc,i) incompressibility flag upkey (int,sc,i) mixed uP key (/=0 for m. nprophy (int,sc,i) number of prophy values
                                         User input invariant set number
С
                                        incompressibility flag (.true. for incompressible)
С
                                       mixed uP key (/=0 for mixed uP formulation)
C
С
       prophy (dp,ar(*),i) material property array nFib (int,sc,i) number of fibers invariants
С
С
С
       invar
                    dp,ar(3)
                                         invariants
       fibDir
                    (dp,ar(3,*),i) original fiber directions array
C
                    (int,sc,i) number of invariants
(dp,ar(*),i) set of Invariants
С
       ninv
С
       Invar
C
С
      output arguments
С
      _____
                                          value of potential
       potential dp,sc
C
       pD1 dp,ar(*)
                                          1st derivatives of potential wrt Invar(*)
C
       pD2
                  dp,ar(*)
                                          2nd derivatives of potential wrt Invar(*)
С
С
       pD3
                  dp,ar(*)
                                           3rd derivatives of potential wrt J
```

2.4.5. Subroutine UserCreep (Defining Creep Material Behavior)

Use the subroutine UserCreep to define creep material behavior. The subroutine is applicable when you issue the **TB** command with the CREEP option, and with TBOPT = 100.

UserCreep supports shared memory and distributed parallel processing; however, you are responsible for ensuring that your code can use parallel processing.

The subroutine is called at all integration points of elements for which the material is defined by this command. The program always uses implicit time integration for this creep option. You can use plasticity options (BISO, BKIN, NLISO, PLASTIC) to define the plastic behavior of materials. Creep and plastic strain are calculated simultaneously when both creep and plasticity are defined for a material.

Using this subroutine, you can specify a "uniaxial" creep law that will be generalized to the multi-axial state by the general time-dependent viscoplastic material formulation implemented in the program. You can use and update internal state variables in the subroutine. The number of state variables must be defined (**TB**,STATE).

Please see the **TB** command description for more information.

```
*deck,usercreep
                                    USERDISTRIB
     SUBROUTINE usercreep (impflg, ldstep, isubst, matId , elemId,
                           kDInPt, kLayer, kSecPt, nstatv, nprop,
    ۶
                           prop , time , dtime , temp , dtemp ,
    &
                           toffst, Ustatev, creqv , pres , seqv ,
                           delcr , dcrda)
    &
      *** primary function ***
C
           Define creep laws when creep table options are
           TB, CREEP with TBOPT=100.
C
           Demonstrate how to implement usercreep subroutine
С
С
            Creep equation is
C
               dotcreq := k0 * seqv ^n * creqv ^m * exp (-b/T)
С
С
               seqv is equivalent effective stress (Von-Mises stress)
C
С
               creqv is equivalent effective creep strain
                    is the temperature
С
C
               k0, m, n, b are materials constants,
           This model corresponds to primary creep function TBOPT = 1
С
C
С
                                                          gal 10.01.1998
c Copyright ANSYS. All Rights Reserved.
C
      input arguments
C
С
     ==========
      impflg (in ,sc
                         ,i)
                                         Explicit/implicit integration
C
                                         flag (currently not used)
С
               (in ,sc
                          ,i)
      ldstep
                                         Current load step
С
С
      isubst
               (in ,sc
                          ,i)
                                         Current sub step
      matId
               (in ,sc
                          ,i)
                                         number of material index
С
      elemId (in ,sc
                         ,i)
                                         Element number
C
С
      kDInPt
               (in ,sc
                         ,i)
                                        Material integration point
С
      kLayer
               (in ,sc
                         ,i)
                                         Layer number
                         ,i)
               (in ,sc
      kSecPt
                                         Section point
C
                                         Number of state variables
С
      nstatv
                (in ,sc
                          ,i)
С
      nprop
               (in ,sc
                          ,i)
                                         size of mat properties array
C
С
      prop
               (dp ,ar(*),i)
                                         mat properties array
                                         This array is passed all the creep
C
```

```
constants defined by command
C
С
                                       TBDATA associated with TB, CREEP
                                       (do not use prop(13), as it is used
C
С
                                       elsewhere)
С
                                       at temperature temp.
      time
                                       Current time
С
      dtime
                                       Current time increment
С
С
      temp
                                       Current temperature
                                       Current temperature increment
C
      dt.emp
С
      toffst
               (dp, sc, i)
                                       temperature offset from absolute zero
               (dp ,sc , i)
                                       equivalent effective stress
С
     seqv
С
      creav
               (dp ,sc , i)
                                       equivalent effective creep strain
С
      pres
               (dp ,sc , i)
                                       hydrostatic pressure stress, -(Sxx+Syy+Szz)/3
                                       note that: constitutive consistency is not accounted for
C
                                       if creep strains are function of pressure
С
С
                                                    / output desc
С
     input output arguments
                                       input desc
     С
С
     Ustatev (dp,ar(*), i/o)
                                       user defined iinternal state variables at
                                       time 't' / 't+dt'.
C
                                       This array will be passed in containing the
C
                                       values of these variables at start of the
C
                                       time increment. They must be updated in this
С
                                       subroutine to their values at the end of
С
                                       time increment, if any of these internal
C
                                       state variables are associated with the
С
С
                                       creep behavior.
C
С
     output arguments
С
     ==========
     delcr (dp ,sc , o)
                                       incremental creep strain
C
     dcrda (dp,ar(*), o)
                                       output array
С
С
                                       dcrda(1) - derivitive of incremental creep
                                                  strain to effective stress
С
                                       dcrda(2) - derivitive of incremental creep
С
                                                  strain to creep strain
C
С
С
     local variables
С
     ==========
С
      c1,c2,c3,c4 (dp, sc, 1)
                                       temporary variables as creep constants
                                       temporary variable
      expt (dp ,sc, 1)
С
                 (dp ,sc, 1)
                                       temporary variable
С
C
  --- parameters
```

2.4.6. Subroutine user_tbelastic (Defining Material Linear Elastic Properties)

Subroutine user_tbelastic provides an interface for defining your own material linear elastic properties (**TB**,ELASTIC). The following topics are available:

```
2.4.6.1. Overview of the user_tbelastic Subroutine
```

2.4.6.2. Data Types Supported by user_tbelastic

2.4.6.3. Table (TB) Command for user_tbelastic

2.4.6.4. User Interface for user tbelastic

2.4.6.5. The user_tbelastic API

2.4.6.6. Usage Example for user_tbelastic

2.4.6.1. Overview of the user_tbelastic Subroutine

The user_tbelastic subroutine can define material linear elastic properties as a function of temperature or coordinates. The subroutine is called at the material integration points of elements for which the definition of material elastic properties is a user option. The material properties defined are based on the material coordinate system of the elements.

You can use the subroutine with most current-technology elements and with most nonlinear material models.

For more information about these material models, see the documentation for the **TB** command in the *Command Reference*.

2.4.6.2. Data Types Supported by user_tbelastic

The user_tbelastic subroutine can define the following types of material property data:

Isotropic elasticity with two constants

Define the Young's modulus (EX) and Poisson's ratio (NUXY) material constants

General orthotropic elasticity with nine constants

Define the normal modulus, shear modulus, and minor Poisson's ratios. The order is as follows: EX, EY, EZ, GXY, GXZ, GYZ, NUXY, NUXZ, NUYZ. All nine constants must be defined; no default values are assigned.

Anisotropic elasticity with 21 constants

Define the material elastic stiffness matrix. The matrix consists of 21 constants, and all must be defined.

2.4.6.3. Table (TB) Command for user_tbelastic

Issue a **TB** command using the following syntax to access the user_tbelastic subroutine interface:

TB, ELASTIC, mat,, npts, USER

The ELASTIC argument accesses the elastic material property data table. (For more information, see the documentation for the **TB** command's ELASTIC option in the *Command Reference*.)

The mat value represents the material number, and the npts value is the number of material constants.

The USER argument accesses the interface to the user_tbelastic subroutine.

2.4.6.4. User Interface for user thelastic

The user_tbelastic interface consists of six arguments, as follows:

 Four input arguments for the element number, material number, coordinate array, and temperature

- One input/output argument for the number of material constants
- One output argument consisting of the material constants array

The syntax is as follows: SUBROUTINE user_tbelastic(elemId, matId, coords, temp, nprop, prop)

Argument	Input (I) or Output (O)	Definition
elemId	I	Element number
matId	I	Material number
coords	I	Coordinates of material integration point at initial configuration (geometry)
temp	I	Current temperature at material integration point
nprop	1/0	Number of constants to be returned (input) or actually returned (output), as follows: 2 - isotropic elasticity 9 - orthotropic elasticity 21 - anisotropic elasticity The value for this argument is obtained via the TB,ELASTIC command, and is passed into the subroutine. However, you can redefine this value in the subroutine, which then returns it.
prop	0	The material elastic constants to be defined

2.4.6.5. The user_tbelastic API

Following is the interface to the user_tbelastic subroutine:

```
*deck,user_tbelastic
                                USERDISTRIB
    SUBROUTINE user_tbelastic(elemId, matId, coords, temp,
                       nprop, prop)
   *** primary function ***
С
         user interface for elastic material constants
С
c Copyright ANSYS. All Rights Reserved.
C
С
    input arguments
С
     elemId (in, sc
                          Element number
                   , i)
С
c {\tt matId} (in, sc , i) Number of material index
```

```
С
      temp
             (dp, sc , i) Current temperature
С
             (dp, ar(5), i)
                             Coordinates at initial configuration
C
      coords
С
                              For continuum elements:
С
                               1-3: coordinates of integration point
                               4-5: not used
C
                              For line elements:
С
С
                               1-3: coordinates of integration point
                                      along line member axis
C
С
                               4-5: offsets in element y and z directions
С
С
     output arguments
С
     ______
     nprop (in, sc , o)
                             Number of constants
С
С
                                2 - isotropic elasticity
                                9 - orthotropic elasticity
С
                                21 - anisotropic elasticity
С
             (dp, ar(*), o)
                             Material elastic constants (stiffness)
С
      prop
С
     local variables
C
С
c --- parameters
С
```

2.4.6.6. Usage Example for user_tbelastic

In this example, three elements in parallel are subjected to uniaxial tension.

Element 1 is a SOLID185 element defined via the MP command with linear isotropic elasticity.

Element 2 is a SOLID185 element defined via the user-defined elastic material properties interface.

Element 3 is a SHELL181 element defined via the user-defined elastic material properties interface.

Solid elements are a unit cubic with a 1 mm edge. The shell element is a unit square with a 1 mm edge. The Young's modulus is 210e3 MPa, and the Poisson's ratio is 0.3.

Example Input

```
/bat.ch
/com
/com example for user elastic material property interface
/com
/com element 1 solid185 defined via standard MP command
/com element 2 solid185 defined using ansys elastic material interface
/com element 3 shell181 defined using ansys elastic material interface
/com
/prep7
esize,,1
et,1,185
et,2,181
mp,ex,1,210e3
mp, nuxy, 1, 0.3
tb,elastic,2,1,2,user ! user-defined elastic material interface
! SOLID185 element
mat,2
block,,1,,1,,1
vmesh,1
```

```
mat,1
block,,1,,1,,1
vmesh.2
! SHELL181 element
sectype, 1, shell
secdata, 0.100000,1
secdata, 0.100000,2
rect,,1,,1
secn,1
mat,2
type,2
amesh,1
elist,all,all
nsel,s,loc,x
d,all,ux
nsel,s,loc,y
d,all,uy
nsel,s,loc,z
d,all,uz
/solu
nsel,s,loc,x,1
d,all,ux,0.05
alls
solve
fini
/post1
set,1
pres,s
pres, epel
fini
```

2.4.7. Subroutine userfc (Defining Your Own Failure Criteria)

```
*deck,userfc
                                 USERDISTRIB
     subroutine userfc (elem, matlay, iott, tem, elim, slim,
                        eps, sig, nfcOut, fc)
    primary function: user subroutine for defining your own failure criterion
c *** secondary functions: none
c *** user programmable functions may not be used in parallel processing ***
     this is currently only available with
C
          *** Copyright ANSYS. All Rights Reserved.
С
          *** ansys, inc.
С
c *** Notice - This file contains ANSYS Confidential information ***
С
  input arguments:
C
С
     variable (typ,siz,intent)
                                description
                                - element number
            (int,sc,in)
     elem
C
С
     elim
              (dp,ar(9),in)
                             - failure strains at the current temperature
                                   (see FC command input with Lab1 = EPEL)
     slim
                                - failure stresses and coupling coefficients
С
              (dp,ar(12),in)
                                   at the current temperature
С
                                    (see FC command input with Lab1 = S)
С
С
      eps
               (dp,ar(6),in)
                                - vector of strains
               (dp,ar(6),in)
                                - vector of stresses
С
     sig
               (dp,sc,in)
                                - temperature at this point in the model
С
     tem
С
     matlay
              (int,sc,in)
                                - material number
С
      iott
               (int,sc,in)
                                - unit number for writing
C
c output arguments:
```

```
c variable (typ,siz,intent) description
c nfcOut (int,sc, out) - number of user fc computed
c fc (dp,ar(9),out) - user failure criterion
c
```

2.4.8. Subroutine userCZM (Defining Your Own Cohesive Zone Material)

Define your own interfacial cohesive material law via the TB,CZM,,,,USER command.

Issue the **TBDATA** command to define the material constants. Data may be temperature-dependent and is interpolated at the current temperature of the material integration point and passed to the subroutine.

For more information, see User-Defined Cohesive Material (UserCZM) and Using State Variables with the UserCZM Subroutine in the *Material Reference*.

Following is the user cohesive material interface:

```
*deck,userCZM
                                USERDISTRIB
     subroutine userCZM (matId, elemId, kMatIntPt, ldstep,isubst,
                        keycut, ncomp, nProp, nstatev,
                        Time, dTime, Temp, dTemp,
    &
    &
                        coords, prop, Strain, dStrain,
                        stress, dsdePl, sedEl, sedPl, statev,
    &
                        var1, var2, var3, var4, var5)
С
C*
     *** primary function ***
С
С
           user cohesive zone model example
С
С
С
            TB,CZM,mat,NTEMP,NPTS,user
               TBTEMP if mat. constants are temperature dependent
C
               TBDATA define material constants
С
С
С
     input arguments
     ==========
С
      matId (int,sc,in)
С
     elemId (int,sc,in)
                                      element #
С
     kMatIntPt (int,sc,in)
                                     material integration point #
С
С
     ldstep (int,sc,in)
                                     load step number
      isubst
               (int,sc,in)
                                      substep number
С
С
      ncomp
               (int,sc,in)
                                      number of stress, strain components
С
      nProp
               (int,sc,in)
                                      Number of material ocnstants
     nstatev (int,sc,in)
                                      Number of state variables
C
С
               (dp ,sc,in)
                                      temperature at beginning of time increment
С
      Temp
               (dp ,sc,in)
                                      temperature increment
      dTemp
С
С
      Time
               (dp ,sc,in)
                                      time at beginning of increment (t)
                                      time increment (dt)
      dTime
               (dp ,sc,in)
С
С
      prop
              (dp,ar(nprop),i)
                                      Material constants defined by TB command
С
      Strain
               (dp,ar(ncomp),i)
                                      Interface separation at beginning of time increment
С
                                      Interface separation increment
С
      dStrain
              (dp,ar(ncomp),i)
С
      coords
              (dp,ar(3),i)
                                       current coordinates
C
С
     output arguments
С
     ______
С
      stress (dp,ar(nTesn),io)
                                      Traction stress
```

```
C
     sedEl
            (dp,sc,io)
                                elastic work
С
     sedPl
            (dp,sc,io)
                                plastic work
     keycut (int,sc,io)
                                loading bisect/cut control
С
                                0 - no bisect/cut
С
С
                                1 - bisect/cut
                                (factor will be determined by ANSYS solution control)
C
    dsdePl
           (dp,ar(ncomp,ncomp),io) consistent tangent jacobian matrix
С
С
   input output arguments
C
   ______
                               user defined solution state variables
    statev (dp,ar(nstatev,io)
С
С
С
    С
                              currently not used
С
    var1, var2, var3, var4, var5
С
С
    local variables
    С
С
    debugflag (in,sc, 1)
                                 debugflag to print debug information
С
C
```

2.4.9. Subroutine userswstrain (Defining Your Own Swelling Laws)

You can define your own swelling strain option via the TB,SWELL,,,,USER command.

Use the **TBDATA** command to define the material constants. Data may be temperature-dependent and is interpolated at the current temperature of the material integration point and passed to the subroutine.

For more information, see Swelling in the *Material Reference*.

```
*deck.userswstrain
                                      USERDISTRIB
     subroutine userswstrain (elemId, kMatPoint
    &,
                              matId, nprop, propv
    &,
                              time,
                                      dtime
    &,
                                      defv
                              efvs.
    &,
                              sweqt, dsweq
    &,
                              swvi,
                                      swvo)
C
c *** primary function:
                         compute user defined swelling strain
c *** Notice - This file contains ANSYS Confidential information ***
c Copyright ANSYS. All Rights Reserved.
С
c input arguments:
    matId (int,sc,in)
                              - material #
C
     elemId (int,sc,in)
                             - element #
С
С
    kMatPoint(int,sc,in)
                              - element integration point #
    nprop (int,sc,in)
                               - number of material constants
С
             (int,sc,in)
                               - current time
С
     time
    dtime (int,sc,in)
                             - current time increment
С
    propv (dp,ar(nprop),in) - array of material constants
С
С
                                 (the data input via TBDATA command)
     efvs (dp,ar(*) ,in) - field variables array
C
                                 efvs(1) - current temperature
efvs(2) - current fluence
С
С
                          ,in) - incremental field variables array
             (dp,ar(*)
    defv
C
С
                                 defv(1) - temperature increment
                                 defv(2) - fluence increment
```

```
c sweqt (dp,sc ,in) - equivalent swelling strain at time t
С
c output arguments:
   dsweq (dp,sc
                     ,in) - incremental equivalent swelling strain
С
c not used arguments:
   swvi (dp,sc
                      ,in) - not currently used
C
                      ,in) - not currently used
С
           (dp,sc
С
C************************
С
```

2.4.10. Subroutine userck (Checking User-Defined Material Data)

```
*deck.userck
                                 USERDISTRIB
     subroutine userck (curmat,ntb,tb)
c *** primary function: check the user-defined material data,
                          input with the TB, user command.
C
c *** secondary functions: none
C
С
         *** Copyright ANSYS. All Rights Reserved.
         *** ansys, inc.
С
c *** Notice - This file contains ANSYS Confidential information ***
С
c input arguments:
    curmat (int,sc,in)
                                - current material number
C
                                - dimension of tb
С
              (int,sc,in)
              (dp,ar(ntb),in) - input table
С
C
C
 output arguments:
С
    none
С
```

2.4.11. Supporting Function egen

The function egen (kcomp,ep,nuxy) (function) combines kcomp strain components (ep) per:

```
*deck,egen
     function egen (kcomp,ep,posn)
c primary function: combines strain components to give an "overall" strain
                      used in creep and plasticity calculations
c secondary functions: none
С
     formulation of overall value is by:
С
                                           2 1 2 2
С
        /1
     / -*((ep - ep ) + (ep - ep ) + (ep - ep ) + -*(ep + ep + ep ))
// 2 1 2 2 3 3 1 2 4 5 6
C
С
С
                                 (1 + posn)
С
c *** Notice - This file contains ANSYS Confidential information ***
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
С
  input arguments:
С
     variable (typ,siz,intent) description
              (int,sc,in) - number of components of strain
     kcomp
C
               (dp,ar(6),in) - the strain components
(dp,sc,in) - poisson's ratio
С
     ep
              (dp,sc,in)
     posn
c output arguments:
```

```
c egen (dp,func,out) - the combined strain value c
```

2.4.12. Subroutine userfld (Update User-Defined Field Variables)

You can create your own field variables using via the **INISTATE** command and supported materials (**TB**). Node-based initial state helps to initialize the user-defined field variables that are then used by the **TB** database to evaluate the material properties at an integration point. userfld allows you to examine the current state at the integration point and to modify the field variables as needed.

For more information, see Understanding Field Variables in the Material Reference.

```
*deck,userfld
                          USERDISTRIB
    subroutine userfld
        ( matId, elemId,
             ldstep, isubst, time, dtime,
   &
   δ.
            kDomIntPt, kLayer, kSectPt,
            nDirect, nShear, nComp, nStatev,
            coords,
   δ.
   &
             Temp,dTemp )
*** primary function ***
С
С
С
        Edit Field Variables During Solution
C
С
C
c Copyright ANSYS. All Rights Reserved.
С
    input arguments
С
    -----
С
С
    matId (int.sc.i)
                              material #
    elemId (int,sc,i)
                               element #
С
С
    ldstep (int,sc,i)
                              load step num
C
    isubst (int,sc,i)
                               substep num
    time (int,sc,d)
dtime (int,sc,d)
С
                               time
                               time inc
С
    kDomIntPt (int,sc,i)
                               "k"th domain integration point
C
    kLayer (int,sc,i)
                              "k"th layer
С
                               "k"th Section point
   kSectPt (int,sc,i)
С
   nDirect (int,sc,in)
                              # of direct components
# of shear components
С
   nShear (int,sc,in)
ncomp (int,sc,in)
С
                              nDirect + nShear
C
    nstatev (int,sc,i)
                              Number of state variables
С
С
    Temp
           (dp,sc,in)
                              temperature at beginning of
                               time increment
C
С
            (dp,sc,in)
                               temperature increment
    coords (dp,ar(3),i)
С
                                current coordinates
C
С
    input output arguments
С
    С
С
    output arguments
С
С
List of Supported Field Variable Types
С
C
    FLD_USER_1_TYPE
С
    FLD_USER_2_TYPE
C
   FLD_USER_3_TYPE
```

```
C FLD_USER_4_TYPE
C FLD_USER_5_TYPE
C FLD_USER_6_TYPE
C FLD_USER_7_TYPE
C FLD_USER_8_TYPE
C FLD_USER_9_TYPE
C FLC_USER_9_TYPE
```

2.4.13. Subroutine userthstrain (Defining Your Own Thermal Strain)

You can define the thermal strain via the TB,CTE,,,,USER command and the userthstrain subroutine.

Issue the **TBDATA** command to define the material constants. Data can be field-dependent, and is interpolated at the current field values of the material integration point and passed to the subroutine.

For more information, see Thermal Expansion in the Material Reference.

```
*deck,userthstrain
                                         USERDISTRIB
     subroutine userthstrain (nprop, propv,
                              ncomp, epth)
С
c *** primary function: compute the thermal strain
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
                                 - number of material constants
     nprop (int,sc,in)
C
С
               (dp,ar(ncomp),in) - array of material constants
                                   e.g. coefficients of thermal
С
                                        expansion in x,y,z order
C
    ncomp (int,sc,in)
                                 - number of strain components
С
C
                                   6 for 3-d elements (x,y,z,xy,yz,xz)
                                   4 for plane elements (x,y,z,xy)
3 for beam elements (x,xy,xz)
С
С
                                   1 for line elements (x)
С
C
c output arguments:
     epth (dp,ar(ncomp),out) - thermal strains
C
С
```

2.5. Subroutines for Customizing Contact Interfacial Behavior

The following subroutines enable you to customize contact interfacial behavior for contact elements (CONTA172, CONTA174, CONTA175, CONTA177, and CONTA178):

- 2.5.1. Subroutine usercnprop (Programming Your Own Contact Properties)
- 2.5.2. Subroutine userfric (Writing Your Own Friction Laws)
- 2.5.3. Subroutine userinter (Writing Your Own Contact Interactions)
- 2.5.4. Subroutine userwear (Writing Your Own Wear Law)

The subroutines enable you to:

- Perform a user-defined operation on real constants (subroutine usercnprop (p. 232))
- Write your own friction laws (subroutine userfric (p. 236))

- Write your own contact interactions (subroutine userinter (p. 238))
- Write your own wear law (subroutine userwear (p. 244))

2.5.1. Subroutine usercnprop (Programming Your Own Contact Properties)

This subroutine applies to the CONTA17x contact elements.

```
*deck,usercnprop
                                       USERDISTRIB
      subroutine usercnprop (ndim,coor,nkeyopt,keyopt,nrl,rlconst,
     x nintIn,intIn,nrealIn,realIn,kupdhis,localr,nuval,nintp,usvr,
     x ncomp, stress, strain0, strain, kstat, mu, kcnprop, cnprop, keyerr)
c *** primary function:
                          Allow users to define their own contact properties
С
                           in real constant table
                          This logic is accessed with real constant defined
C
                           by table name: %_CNPROP%
С
С
                           (e.g. rmod,cid,kcnprop,%_CNPROP%)
c *** Notice - This file contains ANSYS Confidential information ***
С
C
          Copyright ANSYS. All Rights Reserved.
С
          *** ansys, inc.
С
C
С
  input arguments:
      variable (type, sze, intent) description
C
С
С
      elem
               (int,sc,in)
                                    - element number
     intpt
               (int,sc,in)
                                    - element integration point number
C
     ndim
               (int,sc,in)
                                    - number of dimensions of the problem
C
С
                                     = 2 2D
                                      = 3 3D
С
С
     nintp
               (int,sc,in)
                                    - the total number of integration points of
                                     an element to be used with this routine
C
С
     nuval
               (int)
                                    - number of additional state variables per
С
                                      integration point
          note: nuval x nintp = nstv(on nsvr command); cannot exceed 840!
C
С
                                    - integer variables passed in
               (int,ar(*),in)
С
     intTn
      intIn
               (int,ar(*),in)
                                    - integer variables passed in
C
                                      intIn(1) = element number
C
                                      intIn(2) = element integration point number
C
                                      intIn(3) = material reference number
С
С
                                      intIn(4) = element type ID number (absolute value)
                                                 > 0 for CONTA171-CONTA177
С
                                                 < 0 for CONTA178
С
С
                                      intIn(5) = real constant ID number
                                      intIn(6) = associated contact nodal number
C
С
                                      intIn(7) = contact indicator
                                                 0: intersection is found
С
                                                 otherwise: no intersection
C
                                      intIn(8) = target element number
C
                                      intIn(9) = flag for forcing sliding
С
                                                 frictional case
С
С
                                                 0 : not forcing
                                                 1 : forcing (Slip direction is
С
С
                                                     defined through CMROT command)
                                      intIn(10) = 1 first pass through
C
                                                    (1st iteration)
C
С
                                                     (useful for initializing state
С
                                                    variables to a non-zero value)
                                                = 2 first pass through key of
C
С
                                                    a restart
                                                = 3 first pass through key of
C
С
                                                    a rezoning
                                      intIn(11) = current load step number
C
```

```
intIn(12) = current substep number
С
                                      intIn(13) = current equilibrium iteration
С
С
                                                   number
                                      intIn(14) = flag for using unsymmetric
С
С
                                                   matrices (nropt,unsym)
                                                   0 : symmetric
C
С
                                                   1 : unsymmetric
                                      intIn(15) = Linear perturbation flag
С
                                                   0 : a general load step
C
С
                                                   1 : a linear perturbation step
                                      intIn(16) = key to indicate output pass
C
                                                   0 : not a output pass
С
С
                                                   1 : output pass
                                      intIn(17) = key to indicate if history-
C
                                                   dependent variables
С
                                                   (user defined) need to be
С
                                                   updated after the substep has
С
                                                   converged
С
С
                                                   1 : update (converged)
С
                                                   0 : do not update (not converged)
С
                                      intIn(18) = key to indicate transient effects
                                                   1 : transient is active
C
С
                                                   0 : transient is not active
С
                                      intIn(19) = large deformation key [nlgeom cmd]
                                                   1 : on
C
                                                   0 : off
С
                                      intIn(20) = analysis type (derived from
С
C
                                                   antype cmd)
                                                   0 : a static analysis
С
                                                   1 : a buckling analysis
С
                                                   2 : a modal analysis
С
С
                                                   3 : a harmonic analysis
                                                   4 : a transient analysis
C
С
                                                   7 : a substructure analysis
                                                   8 : a spectrum analysis
С
                                      intIn(21) = key for displacement & force
C
С
                                                   convergence
С
                                                   1 : converged
                                                   0 : not converged
С
               (dp,ar(*),in)
С
      realIn
                                    - real variables passed in
                                      realIn(1) = contact element length
С
                                      realIn(2) = contact element depth
С
                                      realIn(3) = area associated with the contact
C
С
                                                   detection point
                                      realIn(4) = pinball radius
С
С
                                      realIn(5) = un-scaled normal penalty stiffness
                                      realIn(6) = time (or frequency for a harmonic
С
С
                                                   analysis) at the beginning of this
С
                                                  load step
                                      realIn(7) = time (or frequency for a harmonic
С
                                                   analysis) at the end of this load step
С
                                      realIn(8) = current time value (or frequency value
С
С
                                                   for a harmonic analysis)
                                      realIn(9) = time increment (or frequency increment
C
                                                   for a harmonic analysis) over this
C
С
                                                   substep
С
                                      realIn(10) = temperature offset from absolute
С
                                                    zero
С
                                      realIn(11) = geometric penetration/gap
С
                                                   (current substep)
                                                   > 0 : gap
С
С
                                                   < 0 : penetration
                                      realIn(12) = time increment scaling factor to
С
                                                   be used for structural transient
С
С
                                                   dynamics
                                    - number of key options
      nkeyopt (int,sc,in)
С
               (int,ar(nkeyopt),in)- array containing key options
С
      keyopt
С
                                      keyopt(1) : Select degree of freedom
                                      keyopt(2) : Contact algorithm
C
                                      ... so on (see ANSYS documentation)
С
      nrl
               (int,sc,in)
                                    - number of real constants
```

```
C
      rlconst (dp,ar(nrl),in)
                                    - array containing real constants
                                      Elements CONTA171 to CONTA177
С
                                      rlconst(1) : R1
C
                                      rlconst(2) : R2
С
С
                                      rlconst(3) : FKN
                                      rlconst(4) : FTOLN
С
                                      ... so on (see ANSYS documentation)
С
                                      Element CONTA178
С
                                      rlconst(1) : FKN
C
C
                                      rlconst(2) : GAP
                                      ... so on (see ANSYS documentation)
С
С
С
      kcnprop (int,sc,in)
                                    - the position of constant in the real set
C
                                      (see ANSYS contact element manual)
С
С
      ncomp
               (int,sc,in)
                                    - number of stress/force component
                                      = 9 for CONTA171-CONTA177
С
                                      = 7 for CONTA178
С
      stress
                                    - stress components at the beginning of
С
               (dp,ar(ncomp),in)
                                      the current iteaation/substep.
C
                                      stress(1) = frictional stress in direction 1
C
                                      stress(2) = frictional stress in direction 2
С
                                                   (3D only)
С
                                      stress(3) = contact normal pressure
C
С
                                      > 0 : compression
                                      < 0 : tension
С
C
                                      the above contact traction must be defined in
                                      a local coordinate system (see localr)
С
                                      stress(4) = heat flux (per area)
С
                                                  flowing into contact
С
                                      stress(5) = heat flux (per area)
C
C
                                                   flowing into target
                                      < 0 heat flowing into a surface
С
С
                                      > 0 heat flowing out of a surface
                                      stress(6) = electrical current density
C
С
                                                   (or pore fluid flux density)
С
                                                  (per area) flowing into contact
                                      stress(7) = electrical current density
С
С
                                                   (or pore fluid flux density)
                                                   (per area) flowing into target
С
                                      > 0 current flowing out of a surface
С
                                      < 0 current flowing into a surface
C
С
                                      stress(8) = diffusion flux density
С
                                                   (per area) flowing into contact
С
                                      stress(9) = diffusion flux density
                                                   (per area) flowing into target
С
С
                                      > 0 flux flowing out of a surface
С
                                      < 0 flux flowing into a surface
      strain0 (dp,ar(ncomp),in) - strain components in the end of the previous
С
                                      substep
С
С
                                      (see strain for each component definition)
      strain (dp,ar(ncomp),in) - current strain components
C
                                      strain(1) = slip increment in direction 1
C
                                      strain(2) = slip increment in direction 2
C
С
                                                 (3D only)
                                      strain(3) = contact normal gap/penetration
С
                                      < 0 : gap
C
                                      > 0 : penetration
С
С
                                      strain(4) = temperature at the contact point
                                                  (from TEMP DOF or temperature load)
С
С
                                      strain(5) = temperature at the target point
                                                   (only from TEMP DOF)
С
                                      strain(6) = voltage (or pore pressure)
C
С
                                                  at the contact point
                                      strain(7) = voltage (or pore pressure)
С
С
                                                  at the target point
С
                                      strain(8) = concentrationat the contact point
                                      strain(9) = concentrationat the target point
C
               (int,sc,in)
                                    - contact status at the end of the previous
С
      kstat
                                      substep
```

```
3 : stick
С
                                      2 : sliding
С
                                      1 : open contact (near)
C
С
                                      0 : open contact (far)
С
      mu
               (dp,sc,in)
                                    - The frictional coef at the end of previous
                                      substep
C
               (dp,ar(6),in)
                                    - Coordinates of the contact detection point
С
                                      coor(1) current x
С
                                      coor(2) current y
C
С
                                      coor(3) current z
                                      coor(4) initial x
С
С
                                      coor(5) initial y
C
                                      coor(6) initial z
                                    - the direction cosines of the local surface
      localr
               (dp,ar(3,3),in)
C
                                      coordinate system at contact detection
С
С
                                      localr(1,1),localr(1,2),localr(1,3) in slip
                                                                           direction 1
С
                                      localr(2,1),localr(2,2),localr(2,3) in slip
С
С
                                                                           direction 2
                                      localr(3,1),localr(3,2),localr(3,3) in normal
С
C
С
С
               (dp,ar(nuval,nintp),inout) - additional state variables from
      usvr
                                      previous equilibrium iteration (saved
C
                                      if the nsvr command is used)
C
      kupdhis (int,sc,in)
                                    - key to indicate if history-dependent
С
С
                                      variables (user defined) need to be
С
                                      updated after the substep has converged
                                      1 : update (converged)
С
                                      0 : do not update (not converged)
С
C
   output arguments:
С
C
      variable (type,sze,intent)
                                     description
С
      cnprop
               (dp,ar(5),out)
                                    - user defined real constant value and
                                      derivatives w.r.t. kcnprop position
C
С
                                      cnprop(1) = user defined real constant value
С
                                      (e.g. kcnprop = 3 for normal contact
                                        stiffness FKN.
С
                                        positive as scaling factor;
С
                                        negative value as the absolute value)
С
                                      cnprop(2) = derivative of the real constant
С
                                                  w.r.t. geometric penetration/gap
C
С
                                      cnprop(3) = derivative of the real constant
С
                                                  w.r.t. contact normal pressure
С
                                      cnprop(4) = derivative of the real constant
                                                  w.r.t. temperature at contact
С
С
                                      cnprop(5) = derivative of the real constant
С
                                                  w.r.t. temperature at target
               (dp,ar(nuval,nintp),inout) - updated additional state variables
C
      usvr
                                      They are passed in as the values at the
С
С
                                      beginning of this substep.
С
                                      They are updated to be the values at the
                                      end of this substep
C
                                      Use NSVR command to size usvr array and
C
С
                                      set nuval to same value as number of
С
                                      variables on NSVR commands
                                      Use userou.F to save these values
C
С
                                      on NMISC record for output purposes.
С
                                      The number of user defined output items on
                                      NMISC should be equal or less than NSTV
C
С
                                      on nsvr command). It cannot exceed 120.
С
       keyerr (int,sc,inout)
                                    - key to indicate if there is any element
C
                                      formulation error.
С
                                      The error could be caused by too
С
С
                                      large incremental step, illegal model.
С
                                      = 0 no error (present value before calling)
                                      = 1 some error happens. ANSYS will
C
С
                                      decide to stop the analysis or cutback
                                      the substep (bi-section) based on other
```

```
c user input and information at higher c level.
```

2.5.2. Subroutine userfric (Writing Your Own Friction Laws)

This subroutine applies to the CONTA17x contact elements.

```
*deck,userfric
                                     USERDISTRIB
      subroutine userfric (elem, mat, intpt, nkeyopt, keyopt, nrl, rlconst,
     x ncomp,npropu,uprop,kfirst,kfsteq,kn,kt,elen,kstat,timval,
     x timinc,tcont,ttarg,toffst,dslip,slip,pres,tau,dt,usvr,
     x fdiss, elener, kupdhis, mu, dtdp)
С
c *** primary function:
                           Allow users to write their own friction laws.
                           This logic is accessed with tb,fric with tbopt=user.
C
                          The below demonstration logic is the same as using
С
                           tb, fric for isotropic Coulomb friction.
                           Other friction laws may require more general
С
                           definition of friction forces.
C
c *** secondary function: demonstrate the use of user-written friction laws
                           in this routine:
С
                           a. update history variables
С
                          b. compute consistent tangent matrix
C
  *** Notice - This file contains ANSYS Confidential information ***
С
C
C
          Copyright ANSYS. All Rights Reserved.
          *** ansys, inc.
С
C
С
  input arguments:
     variable (type,sze,intent)
                                    description
C
С
С
      elem
               (int,sc,in)
                                    - element number (label)
               (int,sc,in)
                                    - material reference number
С
     mat.
С
      intpt
               (int,sc,in)
                                    - element integration point number
     nkeyopt (int,sc,in)
                                    - number of key options
С
C
      kevopt
               (int,ar(nkeyopt),in) - array containing key options
                                      keyopt(1) : Select degree of freedom
С
С
                                      keyopt(2) : Contact algorithm
С
                                      ... so on (see ANSYS documentation)
С
      nrl
               (int,sc,in)
                                    - number of real constants
      rlconst (dp,ar(nrl),in)
С
                                    - array containing real constants
С
                                      Elements CONTA171 to CONTA177
                                      rlconst(1) : R1
С
                                      rlconst(2) : R2
C
С
                                      rlconst(3) : FKN
                                      rlconst(4) : FTOLN
С
                                      ... so on (see ANSYS documentation)
C
                                      Element CONTA178
C
                                      rlconst(1) : FKN
C
С
                                      rlconst(2) : GAP
С
                                      ... so on (see ANSYS documentation)
               (int,sc,in)
                                    - no. of friction stress components (1 or 2)
C
      ncomp
С
               (int,sc,in)
                                    - no. of user-defined friction properties
      npropu
               (dp,ar(npropu),in) - user-defined material properties
С
      uprop
      kfirst
                                    - 1 if first time through, 0 otherwise
C
               (int,sc,in)
                                      (useful for initializing state variables
С
                                      to a non-zero value)
С
                                    - 1 if first equilibrium iteration of a
      kfsteg
               (int,sc,in)
C
С
                                      substep, 0 otherwise
                                    - normal penalty stiffness
С
      kn
               (dp,sc,in)
С
               (dp,sc,in)
                                    - tangential penalty stiffness
С
                                      (an initial guess is provided but
                                       the user must pick a suitable
C
```

```
value that allows minimal tangential
С
С
                                       slip during sticking without
                                       adversely affecting the convergence;
С
                                       a possible choice could be kt=mu*kn).
С
С
                                       For Lagrange multiplier method (keyopt(2)=4
                                       use a small kt (several orders of magnitude
C
                                       smaller than mu*pres).
С
                                    - length of contact element
                (dp,sc,in)
С
      elen
      kstat
                (int,sc,inout)
                                    - contact status
C
C
                                      3 : stick
                                      2 : sliding
С
                                      1 : open contact (near)
С
C
                                      0 : open contact (far)
      timval
                (dp,sc,in)
                                    - current time value
С
      timinc
                (dp,sc,in)
                                    - time increment over this substep
С
                                    - contact surface temperature
С
      tcont
                (dp,sc,in)
                                      (from temperature DOF or temperature load)
С
                (dp,sc,in)
                                    - target surface temperature
С
      ttarg
С
                                      (only from temperature DOF)
      toffst
                (dp,sc,in)
                                    - temperature offset from absolute zero
С
      dslip
                (dp,ar(ncomp),in)
                                    - slip increment (current substep)
C
      slip
                (dp,ar(ncomp),inout)- accumulated slip (previous substep)
С
С
      pres
                (dp,sc,in)
                                     normal pressure/force (current substep)
                                      > 0 : compression
C
                                      < 0 : tension
C
                (dp,ar(ncomp),inout) - frictional stress (previous substep)
С
      tau
                                      Lagrange multiplier contribution is added
С
C
                                      if keyopt(2)=4
                (dp,ar(nuval,nintp),inout) - additional state variables from
С
                                      previous equilibrium iteration (saved
С
                                      if the nsvr command is used)
С
      kupdhis
               (int,sc,in)
                                    - key to indicate if history-dependent
C
                                      variables (user defined) need to be
C
С
                                      updated after the substep has converged
С
                                      1 : update (converged)
                                      0 : do not update (not converged)
C
С
С
   output arguments:
      variable (type,sze,intent)
                                     description
С
С
                                    - updated contact status
С
      kstat
                (int,sc,inout)
                (dp,sc inout)
                                    - updated friction coefficient
С
      mu
      slip
                (dp,ar(ncomp),inout)- updated accumulated slip
C
                (dp,ar(ncomp),inout) - updated frictional stress
С
      t.au
                                     - material tangent modulus
С
      dt
                (dp,ar(5,5),out)
С
                                      rows and columns of dt matrix are
                                      associated to:
С
С
                                      row 1 : frictional stress in direction 1
                                      row 2 : frictional stress in direction 2
С
                                      row 3 : normal pressure
C
                                      row 4 : blank
С
                                      row 5 : blank
С
                                      col 1 : sliding in direction 1
C
                                      col 2 : sliding in direction 2
C
                                      col 3 : normal gap
C
                                      col 4 : temperature at contact
С
                                      col 5 : temperature at targte
С
                                      relevant components to be filled in are:
C
С
                                      dt(1,1): d(tau1)/d(slip1)
С
                                      dt(1,2): d(tau1)/d(slip2)
                                      dt(1,3): d(taul)/d(normal gap)
C
С
                                      dt(2,1): d(tau2)/d(slip1)
                                      dt(2,2): d(tau2)/d(slip2)
С
                                      dt(2,3): d(tau2)/d(normal gap)
C
С
                                      dt(3,3): d(pres)/d(normal gap)
                                      dt(3,3) set to kn internally
С
                                      dt(1,4) : d(taul)/d(tcont)
С
                                      dt(1,5) : d(tau1)/d(ttarg)
С
                                      dt(2,4) : d(tau2)/d(tcont)
C
                                      dt(2,5) : d(tau2)/d(ttarg)
С
               (dp,ar(ncomp),out) - partial derivative of the frictional
```

```
C
                                     stress in direction 1/2 w.r.t. normal
С
                                     pressure used in Lagrange multiplier
                                     method (keyopt(2)=3,4).
C
               (dp,ar(nuval,nintp),inout) - updated additional state variables
С
С
                                     For example, mu value and absolute
                                     accumulated slip could be output as follows:
C
                                     usvr(1,intpt) : mu
С
                                     usvr(2,intpt) : abs. acc. slip in dir1
С
                                     usvr(3,intpt) : abs. acc. slip in dir2
C
C
                                     Use NSVR command to size usvr array and
                                     set nuval to same value as number of
C
С
                                     variables on NSVR commands
C
                                     Use userou.F to save these values
                                     on NMISC record for output purposes.
C
                                     The number of user defined output items on
С
С
                                     NMISC should be equal or less than NSTV
                                     on nsvr command). It cannot exceed 120.
С
С
С
      fdiss
               (dp,sc,out)
                                   - incremental frictional dissipation
                                     per unit area
C
      elener
               (dp,sc,out)
                                   - incremental elastic stored energy
C
                                     per unit area
С
C
   fortran parameters (to be defined by the user):
C
C
     variable (type)
                                   description
С
     nuval
              (int)
                                   - number of additional state variables per
С
                                     integration point
С
     nintp
               (int.)
                                   - \mbox{maximum number of integration points of}
                                     an element to be used with this routine
С
С
                                     (14 is the maximum)
          note: nuval x nintp = nstv(on nsvr command); cannot exceed 840!
C
С
  internal variables:
     variable (type,sze)
                               description
C
С
     dtfac
              (dp,sc)
                               - temporary variable
                               - limit frictional stress
     taulim
              (dp,sc)
С
С
     taueq
               (dp,sc)
                               - equivalent frictional stress
С
     dir1
              (dp,sc)
                               - slip increment direction 1
     dir2
               (dp,sc)
                               - slip increment direction 2
C
С
     dslipeq (dp,sc)
                               - equivalent slip increment
                               - frictional stress 1 from prev substep
С
     oldt1
               (dp,sc)
     oldt2
                               - frictional stress 2 from prev substep
С
               (dp,sc)
               (dp,ar(2))
                               - data array for diagnostic message
С
      err
C
```

2.5.3. Subroutine userinter (Writing Your Own Contact Interactions)

This subroutine applies to the CONTA17x contact elements.

```
*deck,userinter
                                      USERDISTRIB
     subroutine userinter (ndim,coor,nkeyopt,keyopt,nrl,rlconst,
     x npropu,uprop,nintIn,intIn,nrealIn,realIn,kupdhis,localr,
     x nuval, nintp, usvr, ncomp, stress, strain0, strain,
     x kstat, mu, dt, dtdp, kdamp, damp, fdiss, elener, keyerr, keyenv)
C
c *** primary function:
                          Allow users to write their own interaction behavior.
С
                           This logic is accessed with tb, inter with tbopt=user.
c *** secondary function: demonstrate the use of user-written interface laws
                           in this routine:
С
                           a. update history variables
С
                           b. compute consistent tangent matrix
C
c *** Notice - This file contains ANSYS Confidential information ***
C
C
          *** Copyright ANSYS. All Rights Reserved.
C
```

```
*** ansys, inc.
С
   input arguments:
C
С
      variable (type,sze,intent)
                                    description
С
               (int,sc,in)
                                    - element number
С
      elem
      intpt
               (int,sc,in)
                                    - element integration point number
С
                                    - number of dimensions of the problem
      ndim
               (int,sc,in)
С
                                      = 2 2D
C
С
                                      = 3 3D
                                    - the total number of integration points of
      nintp
               (int,sc,in)
С
                                      an element to be used with this routine
С
С
      nuval
               (int)
                                    - number of additional state variables per
                                      integration point
С
          note: nuval x nintp = nstv(on nsvr command); cannot exceed 840!
С
С
      intIn
                                    - integer variables passed in
С
               (int,ar(*),in)
                                      intIn(1) = element number
С
                                      intIn(2) = element integration point number
С
                                      intIn(3) = material reference number
C
                                      intIn(4) = element type ID number (absolute value)
C
                                                  > 0 for CONTA171-CONTA177
C
С
                                                  < 0 for CONTA178
                                      intIn(5) = real constant ID number
С
                                      intIn(6) = associated contact nodal number
C
                                      intIn(7) = contact indicator
С
                                                  0: intersection is found
С
C
                                                  otherwise: no intersection
                                      intIn(8) = target element number
С
                                      intIn(9) = flag for forcing sliding
С
                                                  frictional case
С
                                                  0: - not forcing
C
                                                  1: - forcing (Slip direction is
C
                                                       defined through CMROT command)
С
                                      intIn(10) = 1 first pass through
С
                                                     (1st iteration)
С
С
                                                     (useful for initializing state
С
                                                      variables to a non-zero value)
                                                 = 2 first pass through key of
С
С
                                                     a restart
                                                 = 3 first pass through key of
С
                                                    a rezoning
С
                                      intIn(11) = current load step number
C
С
                                      intIn(12) = current substep number
                                      intIn(13) = current equilibrium iteration number
С
С
                                      intIn(14) = flag for using unsymmetric matrices
                                                   (nropt,unsym)
С
С
                                                   0: - symmetric
С
                                                  1: - unsymmetric
                                      intIn(15) = Linear perturbation flag
С
                                                   0: - a general load step
С
                                                   1: - a linear perturbation step
С
                                      intIn(16) = key to indicate output pass
С
                                                   0: not a output pass
C
                                                   1: output pass
C
С
                                      intIn(17) = key for displacement & force
С
                                                   convergence
                                                   1: converged
C
                                                   0: not converged
С
С
                                      intIn(18) = key to indicate transient effects
                                                   1 : transient is active
С
С
                                                   0 : transient is not active
                                      intIn(19) = large deformation key [nlgeom cmd]
С
                                                   1 : on
C
С
                                                   0 : off
                                      intIn(20) = analysis type (derived from antype)
С
С
                                                   0 : a static analysis
С
                                                   1 : a buckling analysis
                                                   2 : a modal analysis
C
                                                   3 : a harmonic analysis
С
                                                   4 : a transient analysis
```

```
7 : a substructure analysis
С
                                                   8 : a spectrum analysis
С
               (dp,ar(*),in)
                                    - real variables passed in
      realIn
C
С
                                      realIn(1) = contact element length
С
                                      realIn(2) = contact element depth
                                      realIn(3) = area associated with the contact
С
                                                   detection point
С
                                      realIn(4) = pinball radius
С
                                      realIn(5) = unscaled normal penalty stiffness
C
C
                                      realIn(6) = time (or frequency for a harmonic
                                                   analysis) at the beginning of this
C
С
                                                   load step
С
                                      realIn(7) = time (or frequency for a harmonic
                                                   analysis) at the end of this load step
C
                                      realIn(8) = current time value (or frequency value
С
С
                                                   for a harmonic analysis)
                                      realIn(9) = time increment (or frequency increment
С
                                                   for a harmonic analysis) over this
С
С
                                                   substep
                                      realIn(10) = temperature offset from absolute
С
C
                                                    zero
                                      realIn(11) = geometric penetration/gap
C
С
                                                    (current substep)
                                                    > 0 : gap
С
                                                    < 0 : penetration
C
                                      realIn(12) = time increment scaling factor to
С
                                                    be used for structural transient
С
C
                                                    dynamics
                                      realIn(13) = convection coefficient (SFE command)
С
                                      realIn(14) = bulk temp (SFE command)
С
      nkeyopt (int,sc,in)
                                    - number of key options
С
      keyopt
               (int,ar(nkeyopt),in) - array containing key options
C
C
                                      keyopt(1) : Select degree of freedom
                                      keyopt(2) : Contact algorithm
С
                                      ... so on (see ANSYS documentation)
С
               (int.sc.in)
                                    - number of real constants
      nrl
C
С
      rlconst
               (dp,ar(nrl),in)
                                    - array containing real constants
                                      Elements CONTA171 to CONTA177
С
                                      rlconst(1) : R1
С
С
                                      rlconst(2) : R2
                                      rlconst(3) : FKN
С
                                      rlconst(4) : FTOLN
С
                                       ... so on (see ANSYS documentation)
C
С
                                      Element CONTA178
С
                                      rlconst(1) : FKN
С
                                      rlconst(2) : GAP
                                      ... so on (see ANSYS documentation)
C
С
      ncomp
               (int,sc,in)
                                    - number of stress/force component
С
                                      = 9 for CONTA171-CONTA177
                                      = 7 for CONTA178
С
               (dp,ar(ncomp),inout)- stress components (current substep)
С
      stress
С
                                      It is passed in as the stress at the beginning
С
                                      of the current substep. It is updated to be
                                      the stress at the end of this current substep
C
                                      stress(1) = frictional stress in direction 1
C
С
                                      stress(2) = frictional stress in direction 2
С
                                                   (3D only)
                                      stress(3) = contact normal pressure
C
С
                                      > 0 : compression
                                      < 0 : tension
С
                                      the above contact traction must be defined in
С
С
                                      a local coordinate system (see localr)
                                      Lagrange multiplier contribution is added
С
                                      if keyopt(2)=3,4
C
С
                                      stress(4) = heat flux (per area)
                                                   flowing into contact
C
                                      stress(5) = heat flux (per area)
С
С
                                                   flowing into target
                                      < 0 heat flowing into a surface
C
                                      > 0 heat flowing out of a surface
С
C
                                      stress(6) = electrical current density
```

```
(or pore fluid flux density)
С
                                                   (per area) flowing into contact
С
                                      stress(7) = electrical current density
С
                                                   (or pore fluid flux density)
С
С
                                                   (per area) flowing into target
                                      > 0 current flowing out of a surface
С
                                      < 0 current flowing into a surface
С
                                      stress(8) = diffusion flux density
С
                                                   (per area) flowing into contact
C
С
                                      stress(9) = diffusion flux density
                                                   (per area) flowing into target
С
                                      > 0 flux flowing out of a surface
С
C
                                      < 0 flux flowing into a surface
               (dp,ar(ncomp),in)
                                    - strain components in the end of the previous
      strain0
C
С
С
                                      (see strain for each component definition)
      strain
                                    - current strain components
С
               (dp,ar(ncomp),in)
                                      strain(1) = slip increment in direction 1
С
С
                                      strain(2) = slip increment in direction 2
                                                   (3D only)
С
                                      strain(3) = contact normal gap/penetration
C
                                      < 0 : gap
C
С
                                      > 0 : penetration
                                      strain(4) = temperature at the contact point
С
                                                  (from TEMP DOF or temperature load)
C
                                      strain(5) = temperature at the target point
С
                                                  (only from TEMP DOF)
С
C
                                      strain(6) = voltage (or pore pressure)
                                                   at the contact point
С
                                      strain(7) = voltage (or pore pressure)
С
                                                   at the target point
С
С
                                      strain(8) = concentrationat the contact point
C
                                      strain(9) = concentrationat the target point
С
С
      kstat
               (int,sc,inout)
                                    - contact status (current substep)
                                      It is passed in as the status at the
C
С
                                      beginning of the current substep.
                                      It is updated to be the status at the
С
                                      end of the current substep
С
С
                                      3 : stick
                                      2 : sliding
С
                                      1 : open contact (near)
С
                                      0 : open contact (far)
C
                                    - Coordinates of the contact detection point
С
      coor
               (dp,ar(6),in)
                                      coor(1) current x
С
С
                                      coor(2) current y
                                      coor(3) current z
С
С
                                      coor(4) initial x
С
                                      coor(5) initial y
                                      coor(6) initial z
C
      localr
               (dp,ar(3,3),in)
                                    - the direction cosines of the local surface
С
                                      coordinate system at contact detection
С
С
                                      localr(1,1), localr(1,2), localr(1,3) in
                                                             slip direction 1
C
                                      localr(2,1), localr(2,2), localr(2,3) in
C
                                                             slip direction 2
С
                                      localr(3,1), localr(3,2), localr(3,3) in
С
                                                              normal direction
C
С
                                    - number of user-defined interaction properties
С
      npropu
               (int.sc.in)
               (dp,ar(npropu),in) - user-defined material properties
C
      uprop
С
               (dp,ar(nuval,nintp),inout)- additional state variables from
С
      usvr
                                       previous equilibrium iteration (saved
C
С
                                       if the nsvr command is used)
      kupdhis
                                    - key to indicate if history-dependent
               (int,sc,in)
С
                                      variables (user defined) need to be
С
С
                                      updated after the substep has converged
                                      1 : update (converged)
C
                                      0 : do not update (not converged)
С
С
```

```
c output arguments:
      variable (type,sze,intent)
                                  description
C
      kstat.
               (int,sc,inout)
                                  - updated contact status
C
С
      stress
               (dp,ar(ncomp),inout) - updated stress components
С
          (dp,ar(ncomp,ncomp),out)- interface stiffness matrix:
С
                                      dt(i,j) defines the partial derivative of
С
                                      the ith stress component at the current
C
С
                                      substep w.r.t. the jth component of the
                                      relative strain increment array.
С
С
                                      If symmetric solver option used, ANSYS will
C
                                      symmetrize the matrix bu averaging the
                                      off-diagonal terms.
C
                                      rows and columns of dt matrix are
С
                                      associated to:
С
                                      row 1 : frictional stress in direction 1
С
                                      row 2 : frictional stress in direction 2
С
С
                                      row 3 : normal pressure
                                         > 0 : compression
С
                                         < 0 : tension
C
                                      row 4 : heat flux out the contact surface
C
С
                                         < 0 heat flowing into contact
                                         > 0 heat flowing out of target
С
                                      row 5 : heat flux out the target surface
C
                                         < 0 heat flowing into target
С
                                         > 0 heat flowing out of target
С
C
                                      row 6 : electrical current density
С
                                               (or pore prssure)
                                              flowing out the contact surface
С
                                         > 0 current flowing out of contact
С
                                          < 0 current flowing into contact
C
C
                                      row 7 : electrical current density
                                               (or pore prssure)
С
                                         > 0 current flowing out of target
С
                                         < 0 current flowing into target
С
С
                                      row 8 : diffusion flux density
С
                                              flowing out the contact surface
                                         > 0 flux flowing out of contact
С
С
                                          < 0 flux flowing into contact
                                      row 9 : diffusion flux density
С
                                         > 0 flux flowing out of target
С
                                         < 0 flux flowing into target
C
С
                                      \operatorname{col} 1 : sliding in direction 1
                                      col 2 : sliding in direction 2
С
С
                                      col 3 : normal gap
                                         < 0 : gap
С
С
                                          > 0 : penetration
С
                                      col 4 : temperature at the contact surface
                                      col 5 : temperature at the target surface
C
                                      col 6 : voltage at the contact surface
С
                                      col 7 : voltage at the target surface
С
С
                                      col 8 : concentration at the contact surface
                                      col 9 : concentration at the target surface
C
                                      relevant components to be filled in are:
C
С
                                      dt(1,1): d(tau1)/d(slip1)
С
                                      dt(1,2): d(tau1)/d(slip2)
                                      dt(1,3): d(taul)/d(normal gap)
C
С
                                      dt(1,4): d(tau1)/d(tempC)
С
                                      dt(1,5): d(tau1)/d(tempT)
                                      dt(1,6): d(tau1)/d(voltC)
C
С
                                      dt(1,7): d(tau1)/d(voltT)
                                      dt(1,8): d(tau1)/d(concC)
С
                                      dt(1,9): d(tau1)/d(concT)
C
С
                                      dt(2,1): d(tau2)/d(slip1)
                                      dt(2,2): d(tau2)/d(slip2)
C
С
                                      dt(2,3): d(tau2)/d(normal gap)
С
                                      dt(3,1): d(pres)/d(slip 1)
C
                                      dt(3,2): d(pres)/d(slip 2)
С
C
                                      dt(3,3): d(pres)/d(normal gap)
```

```
С
                                      dt(4,1): d(fluxC)/d(slip 1)
С
                                      dt(4,2): d(fluxC)/d(slip 2)
C
                                      dt(4,3): d(fluxC)/d(normal gap)
С
С
                                      dt(4,4): d(fluxC)/d(tempC)
                                      dt(4,5): d(fluxC)/d(tempT)
C
                                      dt(4,6): d(fluxC)/d(voltC)
С
                                      dt(4,7): d(fluxC)/d(voltT)
С
                                      dt(4,8): d(fluxC)/d(concC)
C
С
                                      dt(4,9): d(fluxC)/d(concT)
С
                                      dt(5,4): d(fluxT)/d(tempC)
С
С
                                      dt(5,5): d(fluxT)/d(tempT)
                                      dt(5,6): d(fluxT)/d(voltC)
C
                                      dt(5,7): d(fluxT)/d(voltT)
С
                                      dt(5,8): d(fluxT)/d(concC)
С
                                      dt(5,9): d(fluxT)/d(concT)
С
С
С
                                      dt(6,4): d(eleC)/d(tempC)
                                      dt(6,5): d(eleC)/d(tempT)
С
                                      dt(6,6): d(eleC)/d(voltC)
C
                                      dt(6,7): d(eleC)/d(voltT)
С
С
                                      dt(6,8): d(eleC)/d(concC)
                                      dt(6,9): d(eleC)/d(concT)
С
C
                                      dt(7,4): d(eleT)/d(tempC)
С
С
                                      dt(7,5): d(eleT)/d(tempT)
C
                                      dt(7,6): d(eleT)/d(voltC)
                                      dt(7,7): d(eleT)/d(voltT)
С
                                      dt(7,8): d(eleT)/d(concC)
С
                                      dt(7,9): d(eleT)/d(concT)
С
C
                                      dt(8,4): d(diffC)/d(tempC)
C
                                      dt(8,5): d(diffC)/d(tempT)
С
С
                                      dt(8,6): d(diffC)/d(voltC)
                                      dt(8,7): d(diffC)/d(voltT)
С
С
                                      dt(8,8): d(diffC)/d(concC)
С
                                      dt(8,9): d(diffC)/d(concT)
С
С
                                      dt(9,4): d(diffT)/d(tempC)
                                      dt(9,5): d(diffT)/d(tempT)
С
                                      dt(9,6): d(diffT)/d(voltC)
С
                                      dt(9,7): d(diffT)/d(voltT)
С
С
                                      dt(9,8): d(diffT)/d(concC)
                                      dt(9,9): d(diffT)/d(concT)
С
С
      dtdp
                (dp,ar(ncomp),out) - partial derivative of the frictional stress
                                      in direction 1,2 w.r.t. normal pressure
C
С
                                      used in Lagrange multiplier method
С
                                       (keyopt(2)=3,4).
                (dp,ar(3,3),out)
                                    - interface damping matrix (structure only)
C
      damp
                                       it can be used only in Linear perturbation
С
С
                                      modal analysis or transient analysis or
С
                                      harmonic analysis in frequence domain.
                                      damp(i,j) defines the partial derivative of
С
                                      the ith stress component at the current
C
С
                                      substep w.r.t. the jth component of the
С
                                      strain increment rate array.
                                      rows and columns of dt matrix are
C
С
                                      associated to:
С
                                      row 1 : frictional stress in direction 1
                                      row 2 : frictional stress in direction 2
C
С
                                      row 3 : normal pressure
                                      col 1 : sliding rate in direction 1
С
                                      col 2 : sliding rate in direction 2
C
С
                                      col 3 : normal gap rate
                                    - damping matrix index
      kdamp
                (int.sr.out)
С
С
                                      0 : no damping matrix
С
                                      1 : taking damping matrix into account
                (dp,ar(nuval,nintp),inout) - updated additional state variables
С
      usvr
                                      For example, mu value and absolute/relative
С
                                      accumulated slip could be output as follows:
C
```

```
С
                                      usvr(1,intpt) : mu
                                      usvr(2,intpt) : abs. acc. slip in dir1
С
                                      usvr(3,intpt) : abs. acc. slip in dir2
C
                                      usvr(4,intpt) : acc. slip in dir1
С
С
                                      usvr(5,intpt) : acc. slip in dir2
                                      They are passed in as the values at the
C
                                      beginning of this substep. They are updated
С
С
                                      to be the values at the end of this substep.
                                      Use NSVR command to size usvr array and
C
C
                                      set nuval to same value as number of
                                      variables on NSVR commands
C
С
                                      Use userou.F to save these values
C
                                      on NMISC record for output purposes.
                                      The number of user defined output items on
C
                                      NMISC should be equal or less than NSTV
С
С
                                      on nsvr command). It cannot exceed 120.
С
               (dp,sc,inout)
                                    - The current frictional coefficient
С
С
      fdiss
               (dp,sc,out)
                                    - incremental frictional dissipation
                                      per unit area
C
      elener
               (dp,sc,inout)
                                    - Total elastic stored energy
C
                                      per unit area.
С
                                      Previous converged value is passed in and
С
                                      current total should be the output
C
C
                                    - key to indicate if there is any element
С
      keyerr (int,sc,out)
С
                                      formulation error, like
                                      contact status changes abruptly,
C
С
                                      too much penetration.
С
                                      The error could be caused by too
                                      large incremental step, illegal model.
C
                                      = 0 no error (preset value before calling)
C
C
                                      = 1 some error happens. ANSYS will
                                      decide to stop the analysis or cutback
С
С
                                      the substep (bi-section) based on other
                                      user input and information at higher
C
С
С
      keycnv (int,sc,inout)
                                    - key to flag if this element satisfies
                                      the user defined element convergence
C
С
                                      criterion.
                                      = 1, yes, the criterion is satisfied
С
                                        or don't have any criterion at all
С
                                        it is preset value before calling
C
С
                                      = 0, no, the element doesn't satisfy
С
                                        element convergence criterion. If
С
                                        this is the case, the iteration will
                                        not converge even when both force
C
С
                                        and displacement converge
C
  internal variables:
C
С
      variable (type,sze)
                                description
С
```

2.5.4. Subroutine userwear (Writing Your Own Wear Law)

This subroutine applies to CONTA17x contact elements.

```
*** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
С
c input arguments:
      variable (type,sze,intent)
                                     description
C
С
                                    - number of dimensions of the problem
      ndim
               (int,sc,in)
C
                                      = 2 2D
С
                                      = 3 3D
С
      TotWearOld(dp,ar(ndim),in)
                                    - Total Wear at the contact point at the previous substep
C
С
      strain
               (dp,ar(3),in)
                               - current strain components in contact surface coordinate system
                                      strain(1) = slip increment in direction 1
C
                                      strain(2) = slip increment in direction 2
С
C
                                                   (3D only)
                                      strain(3) = contact normal gap/penetration
С
С
                                      < 0 : gap
                                      > 0 : penetration
С
                                - stress components in contact surface coordinate system
С
      stress
               (dp,ar(3),in)
                                      stress(1) = frictional stress in direction 1
С
С
                                      stress(2) = frictional stress in direction 2
                                                  (3D only)
C
                                      stress(3) = contact normal pressure
C
                                      > 0 : compression
С
                                      < 0 : tension
С
      temperature (dp,sc,in)
                                    - temperature
C
      dtime (dp,sc,in)
                                    - time increment
C
      YieldStress (dp,sc,in)
                                    - Yield stress of underlying element (defined only for Plastic material-see do
С
      nTbprop (int,sc,in)
                                    - Number of TBdata for Tb, Wear per field
С
C
      Tbprop (dp,ar(nTbprop,in)
                                    - TB data for the the Tb, Wear option at the given temperature
                                    - Coordinates of the contact detection point
С
               (dp,ar(6),in)
                                       coor(1) current x
С
                                       coor(2) current y
C
                                       coor(3) current z
С
                                       coor(4) initial x
C
                                       coor(5) initial y
С
                                       coor(6) initial z
С
                                     - contact status (current substep)
      kstat
               (int,sc,in)
С
С
                                       3 : stick
                                       2 : sliding
С
                                       1 : open contact (near)
С
С
                                       0 : open contact (far)
                                    - element number
               (int,sc,in)
С
      elem
      intpt
               (int,sc,in)
                                    - element integration point number
С
      localr
               (dp,ar(3,3),in)
                                    - the direction cosines of the local surface
C
                                      coordinate system at contact detection
С
С
                                      localr(1,1), localr(1,2), localr(1,3) in
С
                                                             slip direction 1
                                      localr(2,1), localr(2,2), localr(2,3) in
С
С
                                                             slip direction 2
С
                                      localr(3,1), localr(3,2), localr(3,3) in
                                                              normal direction
C
      intIn
               (int,ar(*),in)
                                    - integer variables passed in
С
С
                                       intIn(1) = target element number the contact element
С
                                                  is in contat with (or is closest)
                                       intIn(2) = Attached contact element number (if any)
С
                                                  to the target element in passed in intIn(1)
C
                                       intIn(3) = number of additional state variables per
С
С
                                                   integration point (nuval)
                                       intIn(4) = the total number of integration points of
C
                                                  an element to be used with this routine (nintp)
С
                                       intIn(5) = key to indicate if history-dependent
С
                                                  variables (user defined) need to be
С
С
                                                   updated after the substep has converged
                                                  1 : update (converged)
С
                                                  0 : do not update (not converged)
C
      realIn
               (dp,ar(*),in)
                                    - real variables passed in
С
                                      realIn(1) = contact element length
С
                                      realIn(2) = contact element depth
С
С
                                      realIn(3) = area associated with the contact
                                                  detection point
С
С
               (dp,ar(nuval*nintp),inout)- additional state variables from
```

```
C
                                           previous equilibrium iteration (saved
                                           if the nsvr command is used
C
      keyopt (int,ar(*),in) - array containing key options for the element rlconst (dp,ar(*),in) - array containing real constants for the element
C
C
c Output Arguments:
      variable (type,sze,intent)
                                      description
С
С
      WearInc (dp,sc)
                                       - Increment in the Wear (magnitude) - User must define
C
      WearDir (dp,ar(ndim),inout) - Direction cosines in global coordinate system
                                          in which wear increment will be applied- Optional
C
С
                                          default coming in -Contact normal direction
```

2.6. Subroutines for Customizing Loads

The following subroutines modify or monitor existing element loading:

- 2.6.1. Subroutine usrefl (Changing Scalar Fields to User-Defined Values)
- 2.6.2. Subroutine userpr (Changing Element Pressure Information)
- 2.6.3. Subroutine usercy (Changing Element Face Convection Surface Information)
- 2.6.4. Subroutine userfx (Changing Element Face Heat Flux Surface Information)
- 2.6.5. Subroutine userch (Changing Element Face Charge Density Surface Information)
- 2.6.6. Subroutine userfd (Calculating the Complex Load Vector for Frequency Domain Logic)
- 2.6.7. Function userpe (Calculating Rotation Caused by Internal Pressure)
- 2.6.8. Subroutine usrsurf116 (Modifying SURF151 and SURF152 Film Coefficients and Bulk Temperatures)
- 2.6.9. Subroutine User116Cond (Calculating the Conductance Coefficient for FLUID116)
- 2.6.10. Subroutine User116Hf (Calculating the Film Coefficient for FLUID116)
- 2.6.11. Subroutine userPartVelAcc (Calculating Particle Velocities and Accelerations of Ocean Waves)
- 2.6.12. Subroutine userPanelHydFor (Calculating Panel Loads Caused by Ocean Loading)

Activate these subroutines via **USRCAL**.

2.6.1. Subroutine usrefl (Changing Scalar Fields to User-Defined Values)

```
USERDISTRIB
*deck,usrefl
     subroutine usrefl (key,iel,ielc,nnod,nodes,time,defalt,nd,dat)
c *** primary function: change the scalar fields (temperatures, fluences,
                  heat generation, etc.) to what user desires.
c *** secondary functions: none
C
С
         in order to activate this user programmable feature,
         the user must enter the usrcal command.
C
С
          this routine is called at each substep of each load step
С
         for which element or nodal temperatures(etc) are used.
C
С
         it is called for each equilibrium iteration.
C
          the call to get the standard ansys input element or nodal values
         is made just before entering this routine.
C
С
С
         *** Copyright ANSYS. All Rights Reserved.
         *** ansys, inc.
C
c *** Notice - This file contains ANSYS Confidential information ***
```

```
C
      typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
C
  input arguments:
С
      variable (typ,siz,intent)
                                   description
C
С
      kev
               (int,sc,in)
                                 - type of data desired
                                   = 1 temperatures
C
                                   = 2 fluences
С
                                   = 3 heat generation rates
С
                                   = 4 moisture contents
C
С
                                   = 5 magnetic virtual displacements
      iel
                                 - element number
               (int,sc,in)
С
             (int,ar(IELCSZ),in) - array of element type characteristics
С
      ielc
С
      nnod
               (int,sc,in)
                                 - number of nodes
               (int,ar(nnod),in) - list of nodes
      nodes
C
                               - time of current substep
С
     time
               (dp,sc,in)
      defalt
                                 - default value (e.g. tunif)
С
              (dp,sc,in)
      nd
               (int,sc,in)
                                 - size of dat array
С
               (dp,ar(nd),inout) - array of data as normally computed by element
С
      dat
С
                                   as selected by key
C
  output arguments:
      variable (typ,siz,intent)
                                   description
C
С
               (dp,ar(nd),inout) - array of data passed back to element
                                     this data represents values at the end
С
                                     of the load step
C
С
         the input argument dat may be used in one of three ways:
С
             1. it may be simply passed thru
C
             2. it may be used as a flag(e.g. if dat(1) = -3.0, use
С
                                    a certain set of logic)
С
             3. it may be completely ignored and instead defined with new logic
С
```

2.6.2. Subroutine userpr (Changing Element Pressure Information)

```
*deck.userpr
                                  USERDISTRIB
     subroutine userpr (ielc,elem,time,ndat,dat)
c *** primary function:
                           change element pressure information.
          *** Copyright ANSYS. All Rights Reserved.
          *** ansys, inc.
С
С
          in order to activate this user programmable feature,
          the user must enter the 'usrcal, userpr' command.
C
          this routine is called at each substep of each load step for which
С
          pressures are used. it is called for each equilibrium iteration.
С
          it is called once per element.
С
С
          the call to get the standard ansys input pressures is made just before
          entering this routine.
C
С
      input arguments:
         variable (typ,siz,intent)
                                      description
С
               (int,ar(IELCSZ),in) - array of element type characteristics
С
         ielc
                                 - element number for operation.
         elem
                  (int,sc,in)
C
С
         time
                  (dp,sc,in)
                                    - time of current substep
С
         ndat.
                  (int,sc,in)
                                    - number of pressure items for this element
                  (dp,ar(ndat,2),inout) - the element pressure vector
С
         dat
                                        (has input values for each corner
С
                                       of each face)
С
      output arguments:
C
         variable (typ,siz,intent)
                                      description
С
С
                  (dp,ar(ndat,2),inout) - the element pressure vector
                                        (defines input values for each corner
C
                                       of each face)
С
```

2.6.3. Subroutine usercy (Changing Element Face Convection Surface Information)

```
*deck,usercv
                                  USERDISTRIB
      subroutine usercv (elem,ielc,time,nr,u, ndat,hc,tb)
  *** primary function: change element face convection surface info
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
C
С
C
          in order to activate this user programmable feature,
          the user must enter the 'usrcal, usercv' command.
С
С
         the input arguments hc and tb may be used in one of three ways:
С
             1. they may be simply passed thru.
C
             2. they may be used as a flag(e.g. if hc(2) = -3.0, use
С
                                    a certain set of logic).
С
             3. they may be completely ignored.
                                     and instead redefined with new logic
C
С
          this routine is called during each substep of each load step.
C
          it is called for each equilibrium iteration.
С
          it is called once per element. it is called only during the heat
          flow load vector formulation stage, and not during the heat flow
С
          evaluation stage.
С
          the call to get the standard ansys input convection surfaces
С
С
          is made just before entering this routine, so this information is
          available to be modified, if desired.
С
С
          velocity-dependent film coefficients can be computed by inputting the
C
С
          velocity as the input film coefficient or bulk temperature or
С
          by inputting the velocity as a function of location in space.
          routine could then compute the effective film coefficient.
С
C
      input arguments:
С
                                    description
         variable (typ,siz,intent)
С
С
         elem
                  (int,sc,in)
                                    - element number for operation.
                (int,ar(IELCSZ),in) - array of element type characteristics
С
         ielc
                                 - time of current substep
         time
C
                  (dp,sc,in)
                                    - number of nodal temperatures
С
                  (int,sc,in)
С
                                          of the element
                                    - vector of most recent values of the
С
                  (dp,ar(nr),in)
С
                                       temperatures
С
         ndat
                  (int,sc,in)
                                    - number of data points per element
                                       for example, for solid70, ndat = 24 = 6*4
C
С
                                       where 6 = faces per element
                                             4 = corners per face
C
С
         hc
                (dp,ar(ndat),inout) - film coefficients
С
                                        (has input values for each corner
C
                                       of each face)
С
         tb
                (dp,ar(ndat),inout) - bulk temperature
                                       (has input values for each corner
```

```
C
                                        of each face)
С
      output arguments:
С
         variable (typ,siz,intent)
                                       description
C
С
               (dp,ar(ndat),inout) - film coefficients
                                        (defines input values for each corner
С
                                        of each face)
С
                (dp,ar(ndat),inout) - bulk temperature
С
         tb
                                        (defines input values for each corner
C
С
                                        of each face)
C
```

2.6.4. Subroutine userfx (Changing Element Face Heat Flux Surface Information)

```
*deck,userfx
                                  USERDISTRIB
      subroutine userfx (ielc,elem,time,nr,u, ndat,dat)
  *** primary function: change element face heat flux surface info
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
С
С
C
          in order to activate this user programmable feature,
          the user must enter the 'usrcal, userfx' command.
С
С
          this routine is called during each substep of each load step.
C
          it is called for each equilibrium iteration.
С
          it is called once per element. it is called only during the heat
          flow load vector formulation stage, and not during the heat flow
С
С
          evaluation stage.
          the call to get the standard ansys input heat flux surfaces
С
С
          is made just before entering this routine, so this information is
С
          available to be modified, if desired.
C
С
     input arguments:
         variable (typ,siz,intent)
                                      description
С
                (int,ar(IELCSZ),in) - array of element type characteristics
C
         ielc
         elem
                  (int,sc,in)
                                    - element number for operation.
С
С
         time
                  (dp,sc,in)
                                    - time of current substep
                                    - number of nodal temperatures
С
         nr
                  (int,sc,in)
С
                                           of the element
                                    - vector of most recent values of the
                  (dp,ar(nr),in)
С
С
                                       temperatures
                  (int,sc,in)
С
         ndat
                                    - number of data points per element
                                        for example, for solid70, ndat = 24 = 6*4
С
                                        where 6 = faces per element
С
                                              4 = corners per face
С
         dat
                (dp,ar(ndat),inout) - fluxes
C
С
                                        (has input values for each corner
                                        of each face)
С
С
C
      output arguments:
         variable (typ,siz,intent)
                                       description
С
С
         dat (dp,ar(ndat),inout) - fluxes
                                        (defines input values for each corner
С
С
                                        of each face)
С
```

2.6.5. Subroutine userch (Changing Element Face Charge Density Surface Information)

```
*deck,userch
                                 USERDISTRIB
     subroutine userch (ielc,ielem,time,nr,u, ndat,dat)
c *** primary function: change element face charge density surface info
          in order to activate this user programmable feature,
С
С
         the user must enter the usrcal command.
С
С
          this routine is called during each substep of each load step.
         it is called once per element. it is called only during the heat
С
         flow load vector formulation stage, and not during the heat flow
C
         evaluation stage.
         the call to get the standard ansys input charge densities of surfaces
C
С
         is made just before entering this routine, so this information is
         available to be modified, if desired.
С
C
         *** Copyright ANSYS. All Rights Reserved.
С
         *** ansys, inc.
C
C
С
    input arguments:
С
        variable (typ,siz,intent)
                                   description
        ielc (int,ar(IELCSZ),in) - array of element type characteristics
C
С
        ielem (int,sc,in) - element number for operation.
                 (dp,sc,in) - time of current substep
(int,sc,in) - number of nodal temperatures
        time
C
С
        nr
                                         of the element
С
                (dp,ar(nr),in) - vector of most recent values of the
C
С
                                      temperatures
                (int,sc,in) - number of data points per element
        ndat
С
        dat (dp,ar(ndat),inout) - fluxes
C
С
С
     output arguments:
       variable (typ,siz,intent)
                                   description
С
        dat (dp,ar(ndat),inout) - fluxes
С
C
С
         the input argument dat may be used in one of three ways:
            1. they may be simply passed thru.
             2. they may be used as a flag(e.g. if dat(2) = -3.0, use
C
                                   a certain set of logic).
С
            3. they may be completely ignored.
                                    and instead redefined with new logic
С
```

2.6.6. Subroutine userfd (Calculating the Complex Load Vector for Frequency Domain Logic)

```
*deck,userfd
                                   USERDISTRIB
      subroutine userfd (nr,kcbrm,kpfor,ktrsur,isur,
     \verb|xcb,do,doext,aread,alenv,denswat,faclen,conac,fluidt,visc,|\\
     x watbas, watcur, watwav, xyzup, tr, accel, puvel, u, zass,
     x forl, zsc, zsc2, pdyn, holdwv)
c *** primary function: compute complex load vector for frequency domain logic
                          for pipe59
c *** secondary functions: none
      -- accessed with keyopt(12) = 2
C
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
```

```
C
     nr
               (int,sc,in) - matrix size
                                - key for reduced matrices/cable option
С
     kcbrm
               (int,sc,in)
                                - keyopt for hydrodynamic printout
     kpfor
               (int,sc,in)
C
                                - keyopt for surface treatment(unfinished)
     ktrsur
              (int,sc,in)
C
С
     isur
               (int,sc,in)
                                - surface flag
     cb
               (dp,sc,in)
                                - buoyancy coefficient (real constant)
С
     do
               (dp,sc,in)
                                - outside diameter of pipe
С
                                - outside diameter of insulation
С
     doext.
               (dp,sc,in)
                                - area of displaced water
     aread
               (dp,sc,in)
C
С
     alenv
               (dp,sc,in)
                                - length of element
                                - water density
     denswat (dp,sc,in)
С
                                - wetted fraction of pipe
С
     faclen
              (dp,sc,in)
                                - added mass per unit length
С
     conac
               (dp,sc,in)
                                - fluid temperature
     fluidt.
             (dp,sc,in)
C
                                - viscosity
С
     visc
              (dp,sc,in)
              (dp,ar(*),in
С
     watbas
                                - water basic table
     watcur
              (dp,ar(*),in
                                - water current table
С
                                - water wave table
              (dp,ar(*),in
С
                                - updated coordinates
С
     xyzup
               (dp,ar(3,2),in)
                               - local to global transformation matrix
               (dp,ar(3,3),in)
C
     tr
     accel
               (dp,ar(3),in)
                                - acceleration vector
C
     puvel
               (int,sc,in)
                                - index for velocities in u matrix
C
               (dp,ar(nr,5),in
                                - displacements and velocities
С
     u
               (dp,ar(nr,nr),in) - mass matrix
С
      zass
               (dp,ar(12),inout) - force vector in element coordinates
C
     forl
               (dp,ar(nr),inout) - real load vector for frequency domain
С
     zsc
               (dp,ar(nr),inout) - complex load vector for frequency domain
С
     zsc2
C
С
  output arguments:
              (dp,ar(12),inout) - force vector in element coordinates
С
     forl
               (dp,ar(nr),inout) - real load vector for frequency domain
     ZSC
C
     zsc2
               (dp,ar(nr),inout) - complex load vector for frequency domain
С
     pdyn
С
               (dp,ar(2),out)
                                - dynamic pressure
     holdwv
              (dp,ar(60),out)
                               - wave information held for printout
С
C
```

2.6.7. Function userpe (Calculating Rotation Caused by Internal Pressure)

```
USERDISTRIB
*deck.userpe
      function userpe (prs,rvrp,angle,ex,nuxy)
                       calculate the rotation caused by internal pressure
c primary function:
                       on an elbow element
                       This function is only called by el18(pipe18)
C
C
                                                if keyopt(5) = 1
c *** Notice - This file contains ANSYS Confidential information ***
          *** Copyright ANSYS. All Rights Reserved.
С
          *** ansys, inc.
C
С
      typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
   input arguments:
     variable (typ,siz,intent)
                                 description
C
      prs
               (dp,ar(5),in)
                                  - pressure vector
С
С
      rvrp
               (dp,ar(11),in)
                                 - real constants(see elements manual)
                                 - subtended angle
      angle
               (dp,sc,in)
С
                                 Young's modulusPoisson's ratio
С
               (dp,sc,in)
С
      nuxy
               (dp,sc,in)
С
  output arguments:
      variable (typ,siz,intent)
                                  description
С
С
              (dp,sc,out)
                                  - rotation caused by internal pressure on the
                                       elbow element
C
```

2.6.8. Subroutine usrsurf116 (Modifying SURF151 and SURF152 Film Coefficients and Bulk Temperatures)

```
USERDISTRIB
*deck,usrsurf116
      subroutine usrsurf116 (elem,ielc,center,jdim,kaxis,time,nr,u,
                      omeg,ndat,temvel,hc,tb,temfluid,mdot,key)
c *** primary function: change element convection surface info
      for surf151 and/or surf152 based on information from fluid116.
      It is called by el151 and el152.
C
C
          in order to activate this user programmable feature,
          the user must have used fluid116 with keyopt(2) = 1.
С
          Further, surf151 and/or surf152 must have keyopt(5) = 1 or 2
C
          (include extra node). Finally, for this routine to do anything,
C
          key(1) and/or key(2) must be reset in this routine to a
С
          nonzero number. There is no usrcal control over this routine.
С
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
С
c *** Notice - This file contains ANSYS Confidential information ***
C
      input arguments:
С
С
        variable (typ,siz,intent)
                                      description
                                    - element number for operation.
               (int,sc,in)
         elem
C
С
         ielc
               (int,ar(IELCSZ),in) - array of element type characteristics
         center (dp,ar(3),in)
                                    - coordinates of center of surface element
С
С
         jdim (int,sc,in)
                                    - dimensionality key
С
                                          1 = 2d
                                          2 = axisymmetric
C
                                          3 = 3d
C
                                    - axis of rotation (keyopt(3) for el152)
С
         kaxis (int.sc.in)
                                         (see getv116 for definition)
C
         time
                                    - time of current substep
С
                (dp,sc,in)
С
         nr
                (int,sc,in)
                                    - number of nodal temperatures
                                          of the element
C
                (dp,ar(nr),in)
                                    - vector of most recent values of the
С
С
                                       temperatures
С
         omeg
                (dp,sc,in)
                                    - spin real constant (may be from table)
С
         ndat
                (int,sc,in)
                                    - number of data points per element
                (dp,ar(ndat),inout) - film coefficients
C
         hc
С
                                       (has input values for each corner
С
                                       of element)
         tb
                (dp,ar(ndat),in) - bulk temperature
С
                                        (has input values for each corner
С
                                       of element)
С
C
         temfluid (dp,sc,in)
                                    - temp of fluid at surf151/152 centroid
                                    - when using kyop5 = 1 or 2
С
                                    - mass flow rate of fluid when using
C
         mdot (dp,sc,in)
                                    - kyop5 = 2 ( 0 otherwise )
С
С
      output arguments:
С
С
         variable (typ,siz,intent)
                                      description
                                    - user defined bulk temperature in excess of
С
         temvel (dp,sc,out)
                                       fluid node temperature
C
         hc
                (dp,ar(ndat),inout) - film coefficients
С
                                        (defines input values for each corner
С
                                       of element)
C
С
         key
                (int,ar(2),out)
                                    - key if to use this logic
                                       key(1) = 0 = no new film coefficient
C
                                        key(1) = 1 = define new film coefficient
С
С
                                        key(2) = 0 do not use any temvel
                                               = 1 use constant temvel
C
                                               = 2 use bilinear variation
С
С
                                                   of temvel by
                                                   redefining tb array
C
C
          this routine is called during each substep of each load step.
С
          it is called for each equilibrium iteration.
```

```
C
          it is called once per element. it is called only during the heat
          flow load vector formulation stage, and not during the heat flow
С
          evaluation stage.
C
          the call to get the standard ansys input convection surfaces
С
С
          is made just before entering this routine, so this information is
          available to be modified, if desired.
C
С
С
          This routine may be thought of as a specialized version of usercv.
          Indeed, el151 and el152 also call usercy. Either (or both, rarely)
C
          could be used.
С
С
          velocity-dependent film coefficients and bulk temperatures can
С
          be computed by using the velocities and other information from
С
С
          Details of this procedure are:
С
             -- SURF151 or SURF152 are 'pasted' onto the actual solid model.
             -- flow rate is input to or is computed by FLUID116,
С
                  with KEYOPT(2) = 1
С
             -- flow rate may be a function of time
С
             -- the user defines nodes on the FLUID116 network to be the same
C
                  nodes as the 'extra' nodes of SURF151 or SURF152. If more
С
                  than one FLUID116 element is attached to one of these nodes,
С
                  the velocities are averaged.
С
             -- SURF151 or SURF152 calls this routine, indirectly, to compute
С
                  the film coefficient and bulk temperature. This routine,
С
                  in turn, gets the average velocity at the 'extra' node
С
С
                  using 'getv116', as shown below. Other quantities brought
C
                  in by getv116 are also averaged.
```

2.6.9. Subroutine User116Cond (Calculating the Conductance Coefficient for FLUID116)

```
*deck, User116Cond
                                      USERDISTRIB
     subroutine User116Cond(elem,prop,rvr,aleng,re,fric,uptot,uttot,
        bco)
c primary function: compute bc for conductance coefficient for fluid116
c *** Notice - This file contains ANSYS Confidential information ***
С
          *** Copyright ANSYS. All Rights Reserved.
С
          *** ansys, inc.
C
c input arguments:
     elem (int,sc,in)
                                - element number
C
              (dp,ar(4),in)
                                - material property vector
С
     prop
С
                                    order is: dens, visc, kxx, c
     rvr
               (dp,ar(24),in)
                                - real constant vector
C
     aleng
               (dp,sc,in)
                                - element length
С
                                - reynold's number
               (dp,sc,in)
С
     re
С
     fric
              (dp,sc,in)
                                - friction factor
     uptot
              (dp,ar(2),in
                               - nodal pressure values from previous iteration
С
              (dp,ar(4),in
                               - nodal temperature values from prev iteration
C
     uttot
                                - the conductance coefficient from TB, fcon
С
     bco
              (dp,sc,inout)
  output arguments:
С
              (dp,sc,inout)
                                - the desired conductance coefficient
```

2.6.10. Subroutine User 116Hf (Calculating the Film Coefficient for FLUID 116)

*deck,User116Hf USERDISTRIB

```
subroutine User116Hf (elem,prop,rvr,aleng,re,uptot,uttot,hf)
c primary function: compute hf for film coefficient for fluid116
c *** Notice - This file contains ANSYS Confidential information ***
C
          *** Copyright ANSYS. All Rights Reserved.
C
          *** ansys, inc.
С
c input arguments:
     elem (int,sc,in)
                                 - element number
               (dp,ar(4),inout) - material property vector
C
    prop
С
                                      order is: dens, visc, kxx, c
     rvr (dp,ar(18),in) - real constant vector
С
     aleng (dp,sc,in) - element length
re (dp,sc,in) - reynold's number
С
С
     re
    uptot (dp,ar(2),in - nodal pressure values from previous iteration uttot (dp,ar(4),in - nodal temperature values from prevs iteration
С
     uttot (dp,ar(4),in
С
              (dp,sc,inout)
     hf
                                  - the film coefficient from TB, hflm
С
                                  - as a function of temp and velocity
С
c output arguments:
                              - the desired film coefficient
     hf (dp,sc,inout)
```

2.6.11. Subroutine userPartVelAcc (Calculating Particle Velocities and Accelerations of Ocean Waves)

The userPartVelAcc subroutine is the primary component of the API for inputting your own wave and current information. The API supports the hydrodynamic capability available with line elements (such as LINK180, BEAM188, BEAM189, PIPE288, and PIPE289). The userPartVelAcc subroutine works with the following subroutines:

- userPartVelAccSetup (p. 255), which initializes the data for use by userPartVelAcc, and
- userWavHt (p. 256), which calculates the wave height for a user-defined wave.

For your convenience, two I/O service subroutines are called by the userPartVelAcc subroutine: wvhybl (p. 257) and wvargu (p. 257).

```
*deck.userPartVelAcc
                                          USERDISTRIB
     subroutine userPartVelAcc (elemId,domInt,xyzg,doIns,depth,denswat,
                                 ncm, pCur, watcur,
                                 nw, pWav, watwav, timval,
                                 argu,eta,vxyz,axyz,ar,pdynam)
        ---- accessed only if kwav .ge. 101 ----
С
    ***** primary function: compute particle velocities and accelerations
C
                             due to waves and current
C
    **** secondary function: compute dynamic pressures
С
C
С
          *** Copyright ANSYS. All Rights Reserved.
          *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
    elemId (int,sc,in)
domInt (int,sc,in)
С
                               - element id
                               - integration point number
С
    xyzg (dp,ar(3),in) - coordinates of point of interest
C
    doIns (dp,sc,in)
depth (dp,sc,in)
denswat (dp,sc,in)
С
                                 - outside diameter with insulation
                               - water depth
С
                               - water density
C
  ncm (int,sc,in) - number of current measurements
```

```
C
      pCur
               (int,sc,in)
                                 - pointer in current table (= 30 at 12.0)
                                       i.e. first item is at watcur(pCur+1)
C
                                  - water current table
               (dp,ar(*),in)
C
      watcur
                                             ic = current reading number
C
С
                                     watcur( 6) = ncm = number of current measurements
                                     watcur(pCur + (ic-1)*6 + 1) = Z Coor
С
                                     watcur(pCur + (ic-1)*6 + 2) = Velocity
С
                                     watcur(pCur + (ic-1)*6 + 3) = Angle
С
                                     watcur(pCur + (ic-1)*6 + 4) = Temperature
C
С
                                     watcur(pCur + (ic-1)*6 + 5) = Spare
                                     watcur(pCur + (ic-1)*6 + 6) = Spare
С
С
      nw
               (dp,sc,in)
                                  - number of wave components
С
      pWav
               (int,sc,in)
                                  - pointer to wave table (= 30 at 12.0)
               (dp,ar(*),in)
                                  - water wave table
C
      watwav
                                     watwav( 6) = nw = number of wave components
С
С
                                     watwav(11) = KWAVE (kwav)
                                     watwav(12) = THETA
С
                                     watwav(13) = WAVLOC (kpeak)
С
С
                                     watwav(14) = KCRC
                                     watwav(15) = KMF
С
                                     watwav(16) = PRKEY
C
                                             iw = wave number
C
                                     watwav(pWav + (iw-1)*6 + 1) = Wave Height
С
                                     watwav(pWav + (iw-1)*6 + 2) = Period
C
                                     watwav(pWav + (iw-1)*6 + 3) = Phase Shift
C
                                     watwav(pWav + (iw-1)*6 + 4) = Wave Length
С
С
                                     watwav(pWav + (iw-1)*6 + 5) = Spare
C
                                     watwav(pWav + (iw-1)*6 + 6) = Spare
               (dp,sc,in)
                                  - current time value
С
      timval
С
  output arguments:
C
        While the below 7 arguments are output, they can also
С
C
        be used as input, based on other ANSYS input.
                                 - position in wave (radians) (passed out only for output)
               (dp,sc,out)
С
      arqu
                                 - total wave height
С
      eta
               (dp,sc,out)
                                 - particle velocities
               (dp,ar(3),out)
     VXVZ
С
С
     axyz
               (dp,ar(3),out)
                                 - particle accelerations
С
               (dp,sc,out)
                                 - radial particle acceleration
     ar
              (dp,sc,out)
                                 - dynamic pressure head
С
     pdynam
С
С
  local variable
C
     phead
             (dp,sc,out)
                                 - pressure head
C
```

2.6.11.1. Subroutine userPartVelAccSetup (Initializing Data for Use by the userPartVelAcc Subroutine)

This subroutine initializes the data for the userPartVelAcc (p. 254) subroutine.

```
*deck,userPartVelAccSetup
                                               USERDISTRIB
     subroutine userPartVelAccSetup ( kch,ptr_Ocean,
     х
                                       nsize, nsizec, nsizew,
                                       dWork, dWorkC, dWorkW,
    х
                                       rkd, wvmax )
         ---- accessed only if kwav .ge. 101 -----
c *** primary function: set up and checking of user wave (and current) theory
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
c Copyright ANSYS. All Rights Reserved.
С
c input arguments:
С
     kch (int,sc,in)
                                - key for checking or defaulting (not used by PIPE288)
     ptr_Ocean (int,sc,in)
                                - storage offset
C
                                - size of ocean basic data
     nsize (int,sc,in)
```

```
С
      nsizec
                (int,sc,in)
                                  - size of ocean current data
С
      nsizew
                (int,sc,in)
                                   - size of ocean wave data
                (dp,ar(*),inout) - raw ocean basic data (dWork = watbas)
      dWork
C
С
                                      watbas( 6) = nReN = number of Reynold's numbers
С
                                      watbas(11) = DEPTH
                                      watbas(12) = MATOC
C
                                      watbas(13) = KFLOOD
С
                                      watbas(14) = Ci
С
                                      watbas(15) = Cb
C
C
                                            pBas = 30 (at Rev 12.0) (to be added to argument list)
                                              ir = Reynold's number number
C
С
                                      watbas(pBas + (ir-1)*9 + 1) = RE
                                      watbas(pBas + (ir-1)*9 + 2) = CDy
С
                                      watbas(pBas + (ir-1)*9 + 3) = CDz
C
                                      watbas(pBas + (ir-1)*9 + 4) = CT
С
                                      watbas(pBas + (ir-1)*9 + 5) = CMy
С
                                      watbas(pBas + (ir-1)*9 + 6) = CMz
С
                (dp,ar(*),inout) - raw ocean current data (dWorkC = watcur)
      dWorkC
С
С
      dWorkW
                (dp,ar(*),inout) - raw ocean wave     data (dworkW = watwav)
                                        - see userPartVelAcc.F for details for watcur and watwav
C
C
  output arguments:
С
С
      dWork
                (dp,ar(*),inout) - adjusted ocean basic data
                (dp,ar(*),inout) - adjusted ocean current data
(dp,ar(*),inout) - adjusted ocean wave data
      dWorkC
С
      dWorkW
C
                (dp,sc,out)
                                  - value of k*d
С
      rkd
                                  - total wave height
      wvmax
                (dp,sc,out)
```

2.6.11.2. Subroutine userWavHt

The userWavHt subroutine calculates the wave height of a user-defined wave for the user-PartVelAcc (p. 254) subroutine.

```
*deck,userWavHt
                                     USERDISTRIB
     subroutine userWavHt (xyzg,doext,depth,nw,pWav,watwav,timval,
                            eta,etadot)
        ---- accessed only if kwave .ge. 101 -----
С
      *** primary function: calculate wave height for user wave
C
С
                                over point at xyzg of the element
      *** secondary functions: none
C
c *** Notice - This file contains ANSYS Confidential information ***
c Copyright ANSYS. All Rights Reserved.
С
С
  input arguments:
              (dp,ar(3),in)
                                   - updated coordinates of point of interest in
C
     xyzq
                                   - outside diameter with insulation
С
      doext
               (dp,sc,in)
С
                                    if timval<0.0, argu = doext
С
     depth
               (dp,sc,in)
                                   - water depth
               (int,sc,in)
                                   - number of waves
C
     vsWa
               (int,sc,in)
                                   - pointer to wave table
C
С
     watwav
              (dp,ar(*),in)
                                   - water wave table
     timval
             (dp,sc,in)
                                   - current time value
С
                                     if timval < 0.0
C
С
                                        pass directly in doext position
С
                                        (used for stream function only)
C
                                     else compute value in wvargu
С
С
  output arguments:
С
     eta
              (dp,sc,out)
                                   - wave height
С
      etadot
             (dp,sc,out)
                                   - time derivative of wave height
C
```

2.6.11.3. Subroutine wvhybl

The wvhybl subroutine computes the ratio of two hyperbolic functions and is intended for use with wave loading. It is a utility subroutine called by the userPartVelAcc (p. 254) subroutine.

```
*deck, wvhybl
     function wvhybl (kclass,x,y)
c *** primary function: to compute the ratio of two hyperbolic functions,
                          specialized to the needs of wave loading.
С
                          The options are as given with kclass below.
С
                          Further, only positive values of x and y are used
C
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
С
     variable (typ, siz, intent) description
C
С
     kclass (int,sc,in)
                                -0 - \cosh(x)/\cosh(y)
                                -1 - \sinh(x)/\cosh(y)
C
                                 -2 - \cosh(x)/\sinh(y)
С
                                -3 - \sinh(x)/\sinh(y)
С
                                - argument of numerator
               (dp,sc,in)
C
     х
               (dp,sc,in)
                                - argument of denominator
С
С
c output arguments:
С
      variable (typ, siz, intent) description
      wvhybl (dp,sc,out) - resulting fraction
С
С
```

2.6.11.4. Subroutine wvargu

The wvargu subroutine computes the appropriate position with regard to the wave. It is a utility subroutine called by the userPartVelAcc (p. 254) subroutine.

```
*deck,wvargu
      function wvargu (kpeak,kmf,wavdat,timval,r,doext)
С
      *** primary function: to find appropriate position wrt wave
      *** secondary functions: none
C
С
c *** Notice - This file contains ANSYS Confidential information ***
C
  input arguments:
С
С
     kpeak
              (int,sc,in)
                                - keyopt for when peak effect occurs
     kmf
               (int,sc,in)
                                - key for maccamy-fuchs adjustment
С
С
     wavdat (dp,ar(6),in)
                                - wave data (from water wave table)
                                       wavdat(1) = wave height(not used)
С
C
                                       wavdat(2) = period
С
                                       wavdat(3) = phase shift
                                       wavdat(4) = wave length
C
             (dp,sc,in)
                                - current time value
С
     timval
                                - radial location of point of interest
С
     r
               (dp,sc,in)
C
              (dp,sc,in)
                                - effective outside diameter of pipe
     doext
С
С
  output arguments:
C
     wvargu (dp,sc,out)
                                - wave position(as determined by the argument)
С
                                  output in radians
C
```

2.6.12. Subroutine userPanelHydFor (Calculating Panel Loads Caused by Ocean Loading)

The userPanelHydFor subroutine applies loads and other effects onto SURF154 surface elements. This capability is accessed via KEYOPT(8) of SURF154, together with data read in via the userOceanRead (p. 258) subroutine.

```
*deck,userPanelHydFor USERDISTRIB
subroutine userPanelHydFor (kPOcean, elemId, intPnt,
x depth, denswat,
x ncm, pCur, watcur,
x nw , pWav, watwav,
x xyzupp, vn,
x presoc,admsoc)

c ---- accessed only if kwave on the OCDATA command .ge. 101 -----
c primary function: Get pressure loading on panel
c secondary functions: Get hydrodynamic mass on panel
c load is applied on SURF154 with keyopt(8)

c *** Notice - This file contains ANSYS Confidential information ***
c Copyright ANSYS. All Rights Reserved.

c parameter definition include files:
```

2.6.12.1. Subroutine userOceanRead

The userOceanRead subroutine reads in ocean data to be used by the userPanelHyd-For (p. 258) subroutine.

```
*deck,userOceanRead
                                       USERDISTRIB
     subroutine userOceanRead (iott,kpr,fUnitNo,iOption,
                                pdWaveData,lenWavDat)
        ---- accessed only if kwave on the OCDATA command .ge. 101 ----
c Primary Function: read in ocean file for later use
c Secondary Functions:
C
c --
c Notice:
C ----
  This routine contains ANSYS, Inc. confidential information
c Copyright ANSYS. All Rights Reserved.
C -----
c input arguments:
    iott (int,sc,in) output unit number, based on then /OUT command
С
                (log,sc,in) print flag, based on the /NOPR command
С
     kpr
     fUnitNo
                (int,sc,in) file unit number, based on the command
С
С
                                 OCREAD, file, ext, dir
С
     iOption
                 (int,sc,in)
                                 integer from the command line, based on
С
                                 OCREAD, file, ext, dir, iOption
     pdWaveData (ptr,sc,out)
                                 pointer to wave data array
C
С
     lenWavDat (int,sc,out)
                                length of wave data
                                 0 = an error, no wave data is stored
```

2.7. Subroutines for Sharing Data Between User Routines

In Mechanical APDL running under Windows, each user routine is built into a separate dynamic link library (DLL). To share data, functions and data must be explicitly exported and imported. The following subroutines enable you to share data between user routines via common-block variables:

- 2.7.1. Subroutine userdata (Store Common Block Functionality and Data)
- 2.7.2. Subroutine usercm.inc (Add Common Block Variables)

For more information, see Sharing Data Between User Routines (p. 137).

2.7.1. Subroutine userdata (Store Common Block Functionality and Data)

```
*deck,userdata
                                    USERDISTRIB
      function getusercmvals(iloc,sz,outdata)
c!DEC$ ATTRIBUTES DLLEXPORT :: getusercmvals
#include "usercm.inc"
                     iloc,sz,getusercmvals
      double precision outdata(*)
      if ( iloc.lt.1.or.iloc+sz.gt.userdatsz) then
          getusercmvals = 0
      else
          outdata(1:sz) = userdata(iloc:iloc+sz)
          getusercmvals = 1
      endif
      return
      end
      function setusercmvals(iloc,sz,indata)
c!DEC$ ATTRIBUTES DLLEXPORT :: setusercmvals
#include "usercm.inc"
      integer
                      iloc,sz,setusercmvals
     double precision indata(*)
     if ( iloc.lt.1.or.iloc+sz.gt.userdatsz) then
          setusercmvals = 0
      else
          userdata(iloc:iloc+sz) = indata(1:sz)
          setusercmvals = 1
      endif
      return
      end
      subroutine initusercmvals(arraysz)
c!DEC$ ATTRIBUTES DLLEXPORT :: initusercmvals
#include "impcom.inc"
#include "ansysdef.inc"
#include "usercm.inc"
      integer arraysz
      external fAnsMemAlloc
      PTRFTN fAnsMemAlloc
      character*16 memlabel
     memlabel = 'userdat'
      userdatptr = fAnsMemAlloc(arraysz,MEM_DOUBLE,memlabel)
     userdatsz = arraysz
     return
      subroutine freeusercmvals()
c!DEC$ ATTRIBUTES DLLEXPORT :: freeusercmvals
#include "usercm.inc"
      external fAnsMemFree
      call fAnsMemFree(userdatptr)
      userdatptr = 0
      return
```

```
function getusercmvalsz()
c!DEC$ ATTRIBUTES DLLEXPORT :: getusercmvalsz
#include "usercm.inc"
   integer getusercmvalsz
   getusercmvalsz = userdatsz
   return
   end
```

2.7.2. Subroutine usercm.inc (Add Common Block Variables)

Use this subroutine with userdata to add more common blocks.

2.8. Running Mechanical APDL as a Subroutine

To call the Mechanical APDL program, use the following:

```
program ansys
```

For multiple calls to subroutine ansys, you must open and close standard input in the calling subroutine. (Usually, input and output are FORTRAN units 5 and 6, respectively.) The calling subroutine cannot use the database access subroutines; however, other user-programmable features can use the database access subroutines freely.

There may be times when Mechanical APDL exits abnormally. Check the file.err file to see if Mechanical APDL wrote an exit code to the file before ending. These error codes may help you to understand what caused the abnormal program exit:

Table	2	.1:	ANS	YS	Exit	Cod	es

Code	Explanation	Code	Explanation
0	Normal Exit	14	XOX Error
1	Stack Error	15	Fatal Error
2	Stack Error	16	Possible Full Disk
3	Stack Error	17	Possible Corrupted or Missing File
4	Stack Error	18	Possible Corrupted DB File
5	Command Line Argument Error	21	Authorized Code Section Entered
6	Accounting File Error	25	Unable to Open X11 Server
7	Auth File Verification Error	30	Quit Signal

Code	Explanation	Code	Explanation
8	Error in Mechanical APDL or End-of-run	31	Failure to Get Signal
11	User Routine Error	>32	System-dependent Error
12	Macro STOP Command		

2.9. Defining Your Own Commands

ANSYS, Inc. provides a set of user subroutines named user01 through user10 for defining custom commands:

- 2.9.1. Function user01
- 2.9.2. Function user02 (Demonstrates Offsetting Selected Nodes)
- 2.9.3. Function user03 (Demonstrates Using Memory)
- 2.9.4. Function user04
- 2.9.5. Functions user05 through user10

To define a custom command:

- 1. Insert the code for the functions you want to perform into subroutine user01 (or user02, etc.).
- 2. Link the subroutine into the program.
- 3. Issue the command /UCMD to define a name for a custom command that calls and executes your subroutine. Use the command format shown below:

/UCMD, Cmd, SRNUM

Cmd

The name for your new command. It can contain any number of characters, but only the first four are significant. The name you specify can not conflict with the name of any command or the names of any other commands or macros.

SRNUM

The number of the subroutine your command should call; that is, a value between 01 and 10. For example, suppose that you create and link in a user subroutine for a parabolic distribution of pressure, and you name that subroutine user01. Issuing the command shown below creates a new command, PARB, that when issued calls your parabolic pressure distribution subroutine:

/UCMD,PARB,1

To make these "custom command" subroutines available in all your sessions, include the /UCMD commands in your start-up file (START.ANS).

You also can use /UCMD to remove a custom command. To do so, simply use a blank value for Cmd, as shown below:

/UCMD,,1

This command removes the PARB command. To list all user-defined command names, issue the command /UCMD.STAT.

2.9.1. Function user01

```
*deck,user01
                                USERDISTRIB
     function user01()
c *** primary function:
                         user routine number 01
C
         *** Copyright ANSYS. All Rights Reserved.
         *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
  /**********************
С
   | this is a user routine that may be used by users to include their
С
    special coding. accesss to this routine is by the command usrl.
   usr1 may be changed by the user using the command /ucmd. the
   | user may then use this routine to call his/her special routines.
  ansys routines to access information in the ansys database may be
С
   | found in the "ansys programmer's manual", available from ansys,inc
C
    see user02 for a simple example usage.
   routines user03 to user10 are also available.
c input arguments: none
  output arguments:
     user01 (int,sc,out)
                              - result code (should be zero)
C
                                  (which is ignored for now)
C
C
С
    Functions for accessing data on the command line
    integer function intinfun(iField) - gets an integer from field iField
C
   double precision function dpinfun(iField) - gets double precision
C
   character*4 ch4infun(iField) - gets (upper case) 4 characters
    character*8 ch8infun(iField) - gets (mixed case) 8 characters
    character*32 ch32infun(iField) - gets (mixed case) 32 characters
С
С
C
#include "impcom.inc"
#include "ansysdef.inc"
      external wringr
     integer wringr
     integer user01, iott
     iott = wrinqr(2)
          ***** USER'S CODE IS INSERTED HERE *****
C
     write (iott,2000)
 2000 format (//' ***** CALL TO ANSYS,INC DUMMY USER01 *****'//)
          ***** do not return this result code in a real user routine
С
     user01 = -654321
       ***** instead return a zero *****
C
     user01 = 0
     return
      end
```

2.9.2. Function user02 (Demonstrates Offsetting Selected Nodes)

*deck,user02	USERDISTRIB	

```
function user02()
c *** primary function:
                        user routine number 02
     --- This demonstration offsets selected nodes with the command:
C
             usr2,dx,dy,dz
C
         *** Copyright ANSYS. All Rights Reserved.
C
         *** ansys, inc.
С
c *** Notice - This file contains ANSYS Confidential information ***
 /**********************
  | see user01 for additional information on user routines
C
   С
c input arguments: none
c output arguments:
     user02 (int,sc,out)
                              - result code (should be zero)
C
                                 (which is ignored for now)
С
    *******************
C
   Functions for accessing data on the command line
C
    integer function intinfun(iField) - gets an integer from field iField
С
    double precision function dpinfun(iField) - gets double precision
С
   character*4 ch4infun(iField) - gets (upper case) 4 characters
character*8 ch8infun(iField) - gets (mixed case) 8 characters
С
C
    character*32 ch32infun(iField) - gets (mixed case) 32 characters
С
    *****************
#include "impcom.inc"
#include "ansysdef.inc"
     external wringr,ndingr,ndgxyz,ndpxyz,erhandler, dpinfun
     integer wringr,ndingr,ndgxyz
     double precision dpinfun
     integer user02, iott, maxnp, i ,ksel
     double precision xyz(3), offset(3)
     maxnp = ndinqr(0,DB_MAXDEFINED)
          ***** get the desired offsets from the command line *****
C
     offset(1) = dpinfun(2)
     offset(2) = dpinfun(3)
     offset(3) = dpinfun(4)
     do i = 1, maxnp
         ksel = ndgxyz (i,xyz(1))
         if (ksel .eq. 1) then
           xyz(1) = xyz(1) + offset(1)
            xyz(2) = xyz(2) + offset(2)
            xyz(3) = xyz(3) + offset(3)
            call ndpxyz (i,xyz(1))
         endif
     enddo
         ***** write to output file *****
     iott = wringr(WR_OUTPUT)
     write (iott,2000)
 2000 format (/' NODE OFFSET COMPLETE '/)
         ***** write to GUI window *****
     call erhandler ('user02',3000,
                2, 'NODE OFFSET COMPLETE', 0.0d0,'')
          ***** required return value *****
С
     user02 = 0
     return
     end
```

2.9.3. Function user03 (Demonstrates Using Memory)

```
*deck,user03
                                USERDISTRIB
    function user03()
c *** primary function:
                         user routine number 03. Gives example of
                         ANSYS Memory usage
         *** Copyright ANSYS. All Rights Reserved.
C
         *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
  /*********************
   see user01 for additional information on user routines
  input arguments: none
  output arguments:
C
     user03 (int,sc,out)
                               - result code (should be zero)
С
                                  (which is ignored for now)
C
С
    Functions for accessing data on the command line
C
    integer function intinfun(iField) - gets an integer from field iField
С
    double precision function dpinfun(iField) - gets double precision
    character*4 ch4infun(iField) - gets (upper case) 4 characters
C
    character*8 ch8infun(iField) - gets (mixed case) 8 characters
    character*32 ch32infun(iField) - gets (mixed case) 32 characters
С
#include "impcom.inc"
#include "ansysdef.inc"
     external wringr, ndingr, ndgxyz, ndnext, fAnsMemAlloc,
               fAnsMemFree, erhandler, parreturn, parstatus
     integer
               wringr, ndingr, ndgxyz, ndnext
     PTRFTN
             fAnsMemAlloc
     integer user03, iott, i, ksel, numnp, node, istat
     double precision xyz(3), xmean, ymean, zmean, stdxyz(3),
               sodx, sody, sodz
c pointers:
     pointer (pdXnodeL,Xnode)
     pointer (pdYnodeL,Ynode)
     pointer (pdZnodeL,Znode)
     double precision Xnode(*), Ynode(*), Znode(*)
     \ensuremath{^{***}} Get nodal xyz locations and calculate standard deviation of
C
     *** x coordinates, y coordinates, & z coordinates
C
     *** get number of currently selected nodes
C
     numnp = ndinqr(0,DB_NUMSELECTED)
     if (numnp .le. 0) go to 999
     *** allocate memory for x, y, & z coordinates of nodes
     pdXnodeL = fAnsMemAlloc(numnp,MEM_DOUBLE,'XCoords ')
     pdYnodeL = fAnsMemAlloc(numnp, MEM_DOUBLE, 'YCoords ')
     pdZnodeL = fAnsMemAlloc(numnp,MEM_DOUBLE,'ZCoords')
 *** loop through all selected nodes
```

```
i = 1
      node = 0
     xmean = 0.0d0
     ymean = 0.0d0
      zmean = 0.0d0
     node = ndnext(node)
      if (node .gt. 0) then
          *** get xyz coordinates
С
          ksel = ndgxyz(node,xyz(1))
          *** store this node's xyz coordinates
C
          Xnode(i) = xyz(1)
          Ynode(i) = xyz(2)
          Znode(i) = xyz(3)
          *** while we're looping, accumulate sums to calculate means
С
          xmean = xmean + xyz(1)
          ymean = ymean + xyz(2)
          zmean = zmean + xyz(3)
          *** increment index
С
          i = i + 1
          *** loop back up for next selected node
С
          goto 10
      endif
      *** node = 0, at the end of node list
С
      *** calculate mean of xyz coordinates
C
      xmean = xmean / numnp
     ymean = ymean / numnp
      zmean = zmean / numnp
      *** calculate standard deviation for xyz coordinates
C
      sodx = 0
      sody = 0
      sodz = 0
      do i = 1, numnp
        sodx = sodx + (Xnode(i) - xmean)**2
        sody = sody + (Ynode(i) - ymean)**2
sodz = sodz + (Znode(i) - zmean)**2
      enddo
      stdxyz(1) = sqrt(sodx / (numnp-1))
      stdxyz(2) = sqrt(sody / (numnp-1))
      stdxyz(3) = sqrt(sodz / (numnp-1))
      ***** write to output file *****
      iott = wrinqr(WR_OUTPUT)
     write (iott,2000) xmean,ymean,zmean,
    х
                        stdxyz(1),stdxyz(2),stdxyz(3)
2000 format (/' MEAN FOR X COORDINATES:',G12.5/
               ' MEAN FOR Y COORDINATES: ',G12.5/
    х
               ' MEAN FOR Z COORDINATES: ',G12.5/
    х
    х
               ' STD FOR X COORDINATES: ',G12.5/
               ' STD FOR Y COORDINATES: ',G12.5/
    x
               ' STD FOR Z COORDINATES: ',G12.5)
     ***** write to GUI window *****
     call erhandler ('user03',5000,2,
    x 'STD FOR X COORDINATES: %G %/
    x STD FOR Y COORDINATES: %G %/
     x STD FOR Z COORDINATES: %G', stdxyz(1),'')
      ***** set _STATUS to 0 for success *****
С
     istat = 0
```

2.9.4. Function user04

```
*deck,user04
                                USERDISTRIB
     function user04()
c *** primary function: user routine number 04; demonstrates gettting a
                        list of nodes attached to a keypoint, line, or area
         *** Copyright ANSYS. All Rights Reserved.
C
         *** ansys, inc.
c *** Notice - This file contains ANSYS Confidential information ***
С
   | see user01 for additional information on user routines
С
c input arguments: none
c output arguments:
     user04 (int,sc,out) - result code (should be zero)
С
С
                                  (which is ignored for now)
    ******************
С
    Functions for accessing data on the command line
С
    integer function intinfun(iField) - gets an integer from field iField
С
    double precision function dpinfun(iField) - gets double precision
    character*4 ch4infun(iField) - gets (upper case) 4 characters
С
С
   character*8 ch8infun(iField) - gets (mixed case) 8 characters
    character*32 ch32infun(iField) - gets (mixed case) 32 characters
C
#include "impcom.inc"
#include "ansysdef.inc"
      external ndkpnt
      external wringr, ndline, ndarea, intinfun
     integer wringr, ndline, ndarea, intinfun
      external
                ch4infun
     character*4 ch4infun
     integer user04, iott, listk(20),listl(20),lista(20), listin(1),
               i, num, ktype, nkpnts, nlines, nareas
     character*4 type, lab2
     iott = wringr (WR_OUTPUT)
c --- setup with: /UCMD,GNSME,4
```

```
C
         !gnsme,group,num,type
С
         ! group = kp, ln, or ar
         ! num = entity number of kp, ln, or ar
C
         ! type = interior, or all
С
С
          ---- see input deck dv-5805s
      lab2 = ch4infun(2)
      write (iott,2010) lab2
 2010 format(/' group name (type of entity) = ',a4)
      num = intinfun(3)
      write (iott,2020) num
 2020 format (' entity number =',i4)
      listin(1) = num
      if (lab2 .ne. 'KP ' ) then
         type = ch4infun(4)
         if (type .eq. 'INTE') then
            write (iott,2030)
 2030
            format (' interior nodes only ')
            ktype = 0
         elseif (type .eq. 'ALL ') then
            write (iott,2040)
 2040
            format (' all (interior and edge/end) nodes ')
            ktype = 1
         else
            write (iott,2050)
 2050
            format ('Only INTE or ALL are acceptable in last field',
            ' on user-written gnsme command')
         endif
      endif
      if (lab2 .eq. 'KP ' ) then
         nkpnts = 0
         call ndkpnt (1,listin(1),nkpnts,listk(1))
         write (iott,2110) nkpnts
         format (' number of nodes on keypoint = ',i4)
         write (iott,2115) (listk(i),i=1,nkpnts)
         format (' node on keypoint = ',i4)
 2115
      elseif (lab2 .eq. 'LN ' ) then
         nlines = ndline (num,ktype,listl(1))
         write (iott,2120) nlines
 2120
         format (' number of nodes on line = ',i4)
         write (iott,2125) (listl(i),i=1,nlines)
 2125
         format (' list of nodes on line'/(3x,i4))
      elseif (lab2 .eq. 'AR ' ) then
         nareas = ndarea (num,ktype,lista(1))
         write (iott,2130) nareas
 2130
         format (' number of nodes on area = ',i4)
         write (iott,2135) (lista(i),i=1,nareas)
 2135
         format (' list of nodes on area'/(3x,i4))
      else
         write (iott,2150)
         format (' Only KP, LN, or AR are acceptable on user-written ',
     x 'gnsme command')
      endif
      user04 = 0
      return
      end
```

2.9.5. Functions user05 through user10

The source code for user subroutines user05, user06, user07, user08, user09, and user10 is identical to function user01 shown above.

2.10. Support Subroutines

The following subroutines are available as a convenience for general applications:

- 2.10.1. Function GetRForce (Getting Nodal Reaction Force Values)
- 2.10.2. Function GetStackDisp (Getting Current Displacement Values)
- 2.10.3. Subroutine ElResultStrt (Getting Load Data from Analysis Results)
- 2.10.4. Subroutine ElResultGet (Getting Results Values at Selected Points)
- 2.10.5. Subroutine Ellnterp (Finding Element Coordinates)

2.10.1. Function GetRForce (Getting Nodal Reaction Force Values)

```
*deck.Get.RForce
     function GetRForce (Node, Label, Value)
c primary function: Get the K * u - F at a node from the rfsum vector.
                       warning: This routine is called after the elements
С
                       are formed, but before solution. Therefore,
                        F is from the current iteration, but
C
С
                        u is from the previous iteration. At convergence,
                        this difference will have little effect.
C
C
                       The computations are done immediately after the
С
                       call to UElMatx.
                Use the RFSUM command to ask for the summation.
С
                 Use *GET, Parm, NODE, num, RF, DOFLAB to access the reaction
C
                         sum from the command line.
C
c secondary functions: Return pointer for fast access
c object/library: usr
c *** Notice - This file contains ANSYS Confidential information ***
C
     Prolog is not CONFIDENTIAL INFORMATION
  input arguments:
     variable (typ,siz,intent) description
C
              (int,sc,in)
С
                                - Node Number (User)
                              - DOF Label (Upper Case)
     Label
              (ch*4,sc,in)
С
                                  'UX ','UY ','TEMP','VOLT','ROTY', etc
C
C
  output arguments:
     GetRForce (int,func,out)
                                - status/pointer
C
                                   = 0 - data not valid
C
                                   > 0 - Rfsum pointer to data for fast access
C
                                         see comments below
C
С
     Value
            (dp,sc,out)
                                 - Solution value for Node, Label
                                   All results are in the nodal coordinate
C
С
                                    system
  example usage:
C
С
        external GetRForce
        integer GetRForce, ptr, Node2
С
       double precision Value
c #include "handlecom.inc" (if Value = Rfsum(ptr) form is to be used)
       ptr = GetRForce (Node2,'UY ',Value)
```

```
c later...
c    Value = Rfsum(ptr)
c    directionID is used to translate label into corresponding position in dislab's position
```

2.10.2. Function GetStackDisp (Getting Current Displacement Values)

```
*deck,GetStackDisp
      function GetStackDisp (Node,Label,Value)
c primary function: Get the displacement at a node from the disp vector
c secondary functions: Return pointer for fast access
c object/library: usr
c *** Notice - This file contains ANSYS Confidential information ***
     Prolog is not CONFIDENTIAL INFORMATION
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
  input arguments:
С
     variable (typ,siz,intent) description
C
     Node (int,sc,in) - Node Number (User)
С
                             - DOF Label (Upper Case)
С
     Label (ch*4,sc,in)
                                 'UX ','UY ','TEMP','VOLT','ROTY', etc
C
 output arguments:
    variable (typ, siz, intent) description
C
     GetStackDisp (int,sc,out) - status/pointer
                                 = 0 - data not valid
С
                                 > 0 - UDisp pointer to data for fast access
С
                                       see comments below
    Value
            (dp,sc,out) - Solution value for Node, Label
C
c example usage:
       external GetStackDisp
c#include "handlecom.inc" (only if UDisp(ptr) form is used
      integer GetStackDisp, ptr, Node2
C
       double precision Value
      ptr = GetStackDisp (Node2,'UY ',Value)
C
    later...
C
       Value = UDisp(ptr)
```

2.10.3. Subroutine ElResultStrt (Getting Load Data from Analysis Results)

```
*deck,ElResultStrt
     subroutine ElResultStrt (Label,Comp,LabAvg,TypeData,nVal,iLoc)
c *** primary function: (post1) Load data for later ElResultGet
c *** Notice - This file contains ANSYS Confidential information ***
               (prolog is not confidential)
c input arguments:
    Label (ch*4,sc,in)
С
                                  - Result Type
C
     Comp
              (ch*4,sc,in)
                                   - Result Component (8 char for ESTR)
С
     LabAvg (ch*4,sc,in)
                                   - 'AVG ' or 'NOAV' ('AVG ' default)
c output arguments:
```

2.10.4. Subroutine ElResultGet (Getting Results Values at Selected Points)

```
*deck,ElResultGet
     subroutine ElResultGet (nPoints, ebest, elcord, TypeData, iLoc,
                            nVal,result)
c *** primary function: (post1) Get results at selected points
c *** Notice - This file contains ANSYS Confidential information ***
               (prolog is not confidential)
c input arguments:
    nPoints (int,sc,in) - Number of evaluation points
                    *** from ElInterp ***
С
     ebest (int,ar(nPoints),in) - Element(s) containing points
elcord (dp,ar(3,nPoints),in) - Element coordinates
С
С
                    *** from ElResultStrt ***
С
                             - Data type code
    TypeData (int,sc,in)
С
    iLoc (int,sc,in)
                                  - Start of selected data
С
    nVal
              (int,sc,in)
                                    - Number of results per point
C
c output arguments:
    Result (dp,ar(nvar,nPoints),out) - Array of results
```

2.10.5. Subroutine ElInterp (Finding Element Coordinates)

```
*deck.ElInterp
     subroutine ElInterp (piFEML, nPoints, xyzPoints, tolInsidein,
                             tolOutsidein,MoveTol,ebest,elcord)
c primary function: Find element numbers containing xyz points
c secondary functions: Find element coordinates of these points
c object/library: upf
c *** Notice - This file contains ANSYS Confidential information ***
     (Prolog is not CONFIDENTIAL INFORMATION)
c input arguments:
  piFEML (ptr,sc,in) - If non 0, pointer of a FEM Object nPoints (int,sc,in) - Number of points to find (do in one group)
c xyzPoints(dp,ar(3,nPoints),in)- XYZ coordinates of each point
 \hbox{$\tt c$} \quad \hbox{tolInsidein}(\hbox{\tt dp,sc,in}) \qquad \quad \hbox{$\tt -$} \; \hbox{Tolerance for point inside element} 
                                         (0.0d0 defaults to 1.0d-4)
С
   tolOutsidein(dp,sc,in)
С
                                    - Maximum distance outside to be associated
                                        with an element (0.0d0 defaults to 0.25)
                                     - Node move tolerance (0.0d0, no move)
c MoveTol (dp,sc,in)
c output arguments:
             (int,ar(nPoints),out) - Best element number for each point
   elcord (dp,ar(3,nPoints),out) - Element coordinates of the point
```

2.11. Access at the Beginning and End of Various Operations

You can access the logic just before a run begins or just after a run ends, and at many other intermediate points, by using the subroutines listed below. These subroutines can perform actions such as evaluating results or performing calculations. (None of the subroutines have input or output arguments.)

Issue the **USRCAL** command (or use an equivalent menu path) to activate or deactivate these subroutines.

User Subroutine	Is Called
UAnBeg [1]	At start-up
USolBeg	Before solution
ULdBeg	Before a load step
USsBeg	Before a substep
UItBeg	Before an iteration
UItFin	After an iteration
USsFin	After a substep
ULdFin	After a load step
USolFin	After solution
UAnFin	At the end of a run

1. The UAnBeg subroutine that allows user access at the start of a run does not require activation by the **USRCAL** command; it is automatically activated when the program is started.

Subroutines USSBeg, UItBeg, UItFin and USSFin default to reading a command macro file from the current working directory whose name is subroutine.mac (that is, ussfin.mac is read by USSFin.F). No user action to relink the program is required for the command macro to be read except that the calling subroutine must be activated by the USRCAL command. The design of the command reading ability of these subroutines is limited to APDL parameter setting commands (*GET, *SET, a = value, etc) and testing for general commands is limited. Commands which are known to work include *DIM, *STATUS. Commands which require another line (*MSG, *VWRITE) are not allowed. Other commands which are known to not work are the solution loading commands (D, F, SFE, and so on). If these capabilities are required, the user will need to create a FORTRAN subroutine and link this subroutine into the program, as described in Understanding User Programmable Features (UPFs) (p. 127).

While parameter substitution into commands is not permitted, USSBeg, and so on were designed to be used in conjunction with dynamic tables and parameter substitution from the user subroutine. As an example, consider a table defined as dS = f(par1), If dS contains values of displacement as a function of PAR1, then dS may be used as a constraint, as

```
*dim,d5,table,10,1,1,PAR1
d5(1)=0,.1,.25,
/solu
d,5,ux,%d5%
```

Modify the value of PAR1 in USSBeg. MAC and the constraint on node 5, ux can then be modified in the middle of a load step.

The following is an example of a valid input that may be read by USSBeg, UItBeg, UItFin and USSFin.

```
/COM, SAMPLE ussfin.mac
a=5
                              ! *get function is ok
b=nx(1)
*get,c,active,solu,Time,cpu ! *get is ok
*dim,array,,6
                               ! array parameters are ok
arrav(1) = 1
array(2) = 2
array(3) = 3
array(4) = 4
array(5) = 5
array(6) = 6
*vleng,3
                                ! vector operations are ok
*vfun,array(4),copy,array(1)
*stat
*stat,array(1)
array(1) =
nnode = ndinqr(0,14)
*dim,array,,nnode
*vget,array(1),NODE,1,NSEL
*stat,array(1)
array(1) =
/eof
/COM, COMMANDS BELOW THIS LINE ARE KNOWN TO NOT WORK
p,1,6,2000
                                ! commands DO NOT work
d,1,uy,.1
*msg,note
THIS IS A TEST MESSAGE
*vwrite,array(1)
(/b = ,f10.4)
```

2.12. Memory-Management Subroutines

The program uses a dynamic memory manager that overlays the system *malloc* and *free* functions and provides a mechanism for accessing the memory from FORTRAN as well as C and C++. The memory manager library for Windows and Linux are:

Windows: The memory manager is in a dynamic linked library, \Program Files | ANSYS Inc\v2020\ANSYS\bin\<platform>\ansMemManager.dll, where <platform> is a directory that uniquely identifies the hardware platform version.

Linux: The memory manager is a shared library in /ansys_inc/v202/ansys/lib/<plat-form>/libansMemManager.so

You may use the system *malloc* and *free* functions or, for FORTRAN, the *allocate* system function. However, you may end up competing with the program for memory, and for large problems there may be insufficient system memory to perform the function.

Alternatively, you can use ANSYS subroutines for memory management.

2.12.1. Using the Memory Manager in a FORTRAN UPF

2.12.2. Using the Memory Manager in a C or C++ UPF

2.12.1. Using the Memory Manager in a FORTRAN UPF

In FORTRAN, dynamic memory is done through *Cray-style* pointers, where a dynamically allocated array is defined via the construct

```
pointer (piArray,Array)
integer Array(*)
```

and memory space for the array is allocated by assigning the pointer, in this case piArray, to the allocated memory space:

```
piArray = fAnsMemAlloc (size,...)
```

or

```
piArray = fAnsMemAllocL (sizeL,...)
```

where size in an integer (4-byte) length variable, and sizeL is a long integer (8-byte) length variable.

To use the memory manager in a FORTRAN UPF, follow these steps:

1. Define the dynamically allocated arrays:

```
pointer (piArray,Array), (pdArray,dArray)
integer Array(*)
double precision dArray(*)
```

2. Initialize the pointers as follows:

```
piArray = PTRFTNNULL
pdArray = PTRFTNNULL
```

3. Allocate space for an array or arrays, as follows:

For integer (4-byte) numbers:

```
piArray = fAnsMemAlloc(ileng,MEM_INTEGER,C16Label)
```

For long integer (8-byte) numbers:

```
piArray = fAnsMemAlloc(ileng,MEM_LONGINT,C16Label)
```

For double-precision numbers:

```
pdArray = fAnsMemAlloc(dleng,MEM_DOUBLE,C16Label)
```

For complex numbers:

```
pcArray = fAnsMemAlloc(cleng,MEM_COMPLEX,C16Label)
```

For real numbers:

```
prArray = fAnsMemAlloc(rleng,MEM_REAL,C16Label)
```

Where the arguments are:

- xleng is the desired size of the array (use fAnsMemAllocL when this size exceeds 2e31 in value)
- MEM_xxx is the keyword indicating the type of data
- C16Label is a character*16 name of the memory block

You must include the ansysdef.inc include file to get the parameter values of MEM_INTEGER, MEM_LONGINT, MEM_DOUBLE, MEM_COMPLEX, and MEM_REAL. The parameter value PTRFTNNULL is defined in improm.inc.

Note:

If there is insufficient memory, fAnsMemAlloc and fAnsMemAllocL return "PTRFTNNULL".

- 4. Use the arrays.
- 5. If necessary, you may either shrink or grow the allocated memory space by using the reallocation routine. In this case, the original pointer and new length must be passed in as follows:

```
piArray = fAnsMemRealloc(piArray,ileng,MEM_INTEGER,C16Label)

Or

piArray = fAnsMemReallocL(piArray,ilengL,MEM_INTEGER,C16Label)
```

6. Deallocate the space using the fAnsMemFree subroutine, as follows:

```
call fAnsMemFree (piArray)
```

The next sections provide Input and output listings for the memory management subroutines in FORTRAN.

For an example using the memory management functions, see: Function user03 (Demonstrates Using Memory) (p. 264) discussed in Defining Your Own Commands (p. 261).

2.12.1.1. Function fAnsMemAlloc (Allocating Space and Returning a Pointer)

```
parallel
                                                                    rlr
deck.fAnsMemAlloc
                        optimize
     function fAnsMemAlloc (iLeng, key, c16Label)
c primary function:
                      Get A Block of Space from mem manager and Return Pointer
c object/library: mem
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
              (int,sc,in)
                                  - length of the block (in data elements)
     iLeng
     c16Label
                 (chr*16,sc,in) - 16 character name for the Block
С
                  (int,sc,in)
                                    - type of data for this block (see ansysdef.inc)
     kev
C
c output arguments:
C
     fAnsMemAlloc (PTRFTN,sc,out)
                                    - Pointer to this data block -- needs to be
                                      tied to a local variable in the calling
```

```
c routine
```

2.12.1.2. Function fAnsMemAllocL (Allocating Space and Returning a Pointer long integer)

```
*deck,fAnsMemAllocL
                         optimize parallel
                                                                       rlr
     function fAnsMemAllocL (iLengL, key, c16Label)
                      Get A Block of Space from mem manager and Return Pointer
c primary function:
c object/library: mem
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
                   (LONG,sc,in)
                                     - length of the block (in data elements)
     iLenqL
                                     - 16 character name for the Block
С
     c16Label
                   (chr*16,sc,in)
                                     - type of data for this block (see ansysdef.inc)
                   (int,sc,in)
С
c output arguments:
С
     fAnsMemAllocL (PTRFTN,sc,out)
                                     - Pointer to this data block -- needs to be
                                       tied to a local variable in the calling
                                       routine
C
```

2.12.1.3. Function fAnsMemRealloc (Reallocating Space and Returning a Pointer)

```
*deck,fAnsMemRealloc
                           optimize parallel
                                                                     rlr
     function fAnsMemRealloc (memPtr, iLeng, key, c16Label)
c primary function:
                      Modify a Block of Space from mem manager and
                      Return Pointer
c object/library: mem
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
                    (PTRFTN,sc,in)
                                     - pointer of block being reallocated
    memPtr
     iLeng
                    (int,sc,in)
(int,sc,in)
                                     - new length of the block (in data elements)
С
С
     key
                                     - type of data needed (see ansysdef.inc)
                    (ch*16,sc,in)
     c16Label
                                     - name for this block
C
c output arguments:
     fAnsMemRealloc (PTRFTN,sc,out)
                                     - Pointer to the new block location
```

2.12.1.4. Functional fAnsMemReallocL (Reallocating Space and Returning a Pointer - long integer)

```
*deck,fAnsMemReallocL optimize parallel rlr
function fAnsMemReallocL (memPtr, iLengL, key, c16Label)

c primary function: Modify a Block of Space from mem manager and c Return Pointer

c object/library: mem

c *** Notice - This file contains ANSYS Confidential information ***
```

```
c input arguments:
c memPtr (PTRFTN,sc,in) - pointer of block being reallocated
c iLengL (LONG,sc,in) - new length of the block (in data elements)
c key (int,sc,in) - type of data needed (see ansysdef.inc)
c c16Label (ch*16,sc,in) - name for this block

c output arguments:
c fAnsMemReallocL (PTRFTN,sc,out) - Pointer to the new block location
```

2.12.1.5. Subroutine fAnsMemFree (Deallocating Space)

```
*deck,fAnsMemFree optimize parallel rlr
subroutine fAnsMemFree (memPtr)

c primary function: Free a Data Block, given a pointer

c object/library: mem

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
    ptr (PTRFTN,sc,inout) - pointer for this block

c output arguments:
    ptr (PTRFTN,sc,inout) - pointer will be set to PTRFTNNULL
```

2.12.2. Using the Memory Manager in a C or C++ UPF

To use the memory manager In C or C++, follow these steps:

1. Define the dynamically allocated arrays:

```
int *iArray;
double *dArray;
```

2. Initialize the pointers as follows:

```
iArray = (int*) NULL;
dArray = (double*) NULL;
```

3. Allocate space for an array or arrays, as follows:

For integer (4-byte) numbers:

```
numBytes = leng*sizeof(int);
iArray = cAnsMemAlloc(numBytes,0,__FILE__,__LINE__);
```

For double-precision numbers:

```
numBytes = leng*sizeof(double);
dArray = cAnsMemAlloc(numBytes,0,__FILE__,_LINE__);
```

Where the arguments are:

- numBytes is the desired amount of memory to allocate (NOTE: this length can be a 64-bit integer)
- The __FILE__ and __LINE__ arguments are used to create a string for naming the memory block

Note:

If there is insufficient memory, cAnsMemAlloc will return a NULL pointer.

- 4. Use the arrays.
- 5. If necessary, you may either shrink or grow the allocated memory space by using the reallocation routine. In this case, the original pointer and new length must be passed in as follows:

```
iArray = cAnsMemRealloc(iArray,newNumBytes,0,__FILE__,_LINE__);
```

6. Deallocate the space using the cAansMemFree subroutine, as follows:

```
call cAnsMemFree (iArray,__FILE__,__LINE__);
```

Note:

the pointer passed into cAnsMemFree is untouched upon leaving this routine.

The next sections provide Input and output listings for the memory management subroutines in C or C++.

2.12.2.1. Function cAnsMemAlloc (Allocating Space and Returning a Pointer)

```
void *cAnsMemAlloc(size_t iLen, INT key, CHAR *cName, INT cLineNum)
^{\star} Allocate a block of memory from the memory manager and return pointer
* PARAMETER LIST:
          - length of block to allocate (in bytes)
   iLen
             - bit patterned descriptive key to be passed to lower level manager
               =0, get memory any way possible
               &MEM_INITIAL, get memory only from initial heap block
               &MEM_GROWTH, get memory only outside of initial heap block
               also stores bit patterns (in bits 17-23) representing various
               allocation groups (solvers, etc)
            - name of calling routine (should be gotten from __FILE__)
   cLineNum - line number of calling routines (gotten from __LINE__)
* RETURN VALUE:
   a pointer to the memory space
*/
```

2.12.2.2. Function cAnsMemRealloc (Reallocating Space and Returning a Pointer)

2.12.2.3. Subroutine cAnsMemFree (Deallocating Space)

```
void cAnsMemFree(void *memPtr, CHAR *cName, INT cLineNum)
/*
   * Free a block of memory, given a pointer
   *
   * PARAMETER LIST:
   * memPtr - pointer to the memory block being deallocated
   * cName - name of calling routine (should be gotten from __FILE__)
   * cLineNum - line number of calling routines (gotten from __LINE__)
   *
   * RETURN VALUE:
   * a pointer to the memory space (will be set to NULL)
   */
```

2.13. Parameter-Processing Subroutines

The product distribution medium contains three subroutines that you can use for parameter processing: pardim, parev1, and pardef.

2.13.1. Subroutine pardim (Creating a Dimensioned Parameter)

```
*deck,pardim
     subroutine pardim (cName,labl4,nDim,nxyz,cLabels)
c *** primary function:
                          create a dimensioned parameter
        *dim,parm32,type,d1,d2,d3,cName1,cName2,cName3
C
        *dim,parm32,type,d1,cName1
С
        *dim,parm32,type,d1,d2,d3,d4,d5,cName1,cName2,cName3,cName4,cName5
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
С
     cName (chr*32,sc,in)
                                 - the name of the parameter to create
               (chr*4,sc,in) - 'TABL' or 'ARRA' or 'CHAR' or 'STRI'
(int,sc,in) - Dimension of array
С
     labl4
    nDim
              (int,sc,in)
C
c nxyz (int,ar(nDim),in) - the dimensions of the array
```

```
c cLabels (chr*32,ar(nDim),in) - Names for the directions in table
c output arguments: none
```

2.13.2. Subroutine parevl (Finding and Evaluating a Parameter)

```
*deck,parevl
     subroutine parevl (ParName, nDim, subc, lvl, dpValue, chValue, kerr)
c *** primary function:
                          find and evaluate a parameter
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
     ParName (chr*(PARMSIZE),sc,in) - the name of the parameter
С
                                        (must be upper case, left justified)
С
                                     - the number of subscripts (0,scaler)
     nDim
               (int.sc.in)
C
С
     subc
               (dp,ar(*),in)
                                     - values for the subscripts (if any)
     1771
               (int,sc,in)
                                     - 0,1 no error output 2, report error
С
С
                                        -1, set kerr flag with no anserr call
  output arguments:
C
                                     - the value of the parameter (may be a
     dpValue (dp,sc,out)
C
С
                                       packed character*8
     chValue (chr*(STRING_MAX_LENG),sc,out) - character output
C
                                      - error flag (0,ok -1,output is packed
С
     kerr
               (int,sc,out)
С
                                        0=ok, 1=error, 2=error but TINY is used
                                         -2, output is string in chValue
C
```

2.13.3. Subroutine pardef (Adding a Parameter)

```
*deck,pardef
     subroutine pardef (cNameIn,ctype,nval,subc,valuein,kerr,string)
c *** primary function:
                          add a parameter to parameter list
c *** Notice - This file contains ANSYS Confidential information ***
С
  input arguments:
C
      cNameIn (chr*(PARMSIZE),sc,in) - name of parameter
C
                              cNameIn is a character variable that
C
                              contains the name of the parameter that
C
                              is to be defined. (Length = PARMSIZE characters)
C
С
                           - 0, dp
                                    1,character 2,string
С
      ctype (int,sc,in)
                              ctype is an integer key which describes
С
                              the type of data that the parameter data
C
                              holds. This would also indicate the
С
                              contents of "value" (arg 5).
С
С
                              0=double precision data
С
                              1=character data packed in value
                              2=character data in string
С
С
      nval (int,sc,in)
                           - number of subscripts
C
С
                              nval is the number of subscripts that the
                              "cNameIn" (arg 1) contains.
С
                              1=single dimensioned variable (ex. x(10))
С
                              2=double dimensioned variable (ex. y(10,3))
C
                              3=triple dimensioned variable (ex. z(10,3,2))
С
                              -1=delete this parameter from the internal
C
С
С
      subc (dp,ar(*),in) - values of subscripts
```

```
C
                              subc is a double precision vector that
С
                              contains the subscripts of "cNameIn" (arg 1).
                              There should be enough values defined to
C
                              match "nval" (arg 3). For example if "x"
C
С
                              was dimensioned as "x(10,3,2)" and you wanted
                              to set "x(5,1,1)=123.0", then "nval" (arg 3)
C
                              should be set to 3, and "subc" should be set
С
                              to 5.0, 1.0, 1.0, and "value" (arg 5) should
С
                              be 123.0. Another example is if "y" was
C
С
                              dimensioned to as "y(20,20)" and you were
                              setting "y(5,8)=987", then "nval" (arg 3) should
С
                              be set to 2 and "subc" should be set to 5.0,
С
С
                              8.0, 0.0, and "value" (arg 5) should be 987.0.
C
                              Remember subroutine "pardef" is only storing
С
С
                              a data value of "cNameIn" or "cNameIn(x,y,z)". The
                              proper dimensions were set by a "*dim" command.
С
С
С
                              Please note that although the values of "subc"
                              should be double precision, subroutine "pardef"
C
                              uses the intrinsic "nint" (nearest integer)
C
                              function to get and use the integer equivalent.
C
С
                              You should also note the "nval" (arg 3) and
С
                             "subc" (arg 4) must fall within the range that was
C
                              set with a "*dim" or "*set" command or an error
С
С
                              will occur.
C
                           - the value for this parameter
С
      valuein(dp,sc,in)
С
                               (should be a packed character*8 if
                              ctype=1. To pack a char into a dp
C
                              variable use "call chtodp(ch8,dp)".
С
C
                              To unpack a dp variable into a char
                              use "call dptoch(dp,ch8)" )
C
С
                              Value is the data value that is to be stored for
                              "cNameIn" (arg 1). If "ctype=1" (arg 2) then this
C
С
                              value would be a "packed character" data from the
С
                              "chtodp" Ansys function.
C
С
  output arguments:
     kerr (int,sc,out) - error flag (0=ok, 1=error)
С
                              kerr is an integer error flag that is
C
                              returned to the calling subroutine. Any
С
С
                              non zero number would indicate an error
                              was detected in subroutine "pardef'
C
  *** mpg pardef < parstore pardim ntableget rdsset<rdmac<rdcmd: define param
C
```

2.14. Other Useful Functions

The program has several miscellaneous functions that you may find useful for working with UPFs:

- The erhandler subroutine displays output messages (notes, warnings, and errors).
- The RunCommand function lets you issue a command from within a user subroutine.
- The GetStackDisp subroutine retrieves current displacement values.
- The /UNDO command writes an "undo" file at critical points as a user subroutine executes.
- The /HOLD command allows you to synchronize multiple tasks.

For further descriptions of erhandler, see Subroutines for Your Convenience (p. 343). For details about the GetStackDisp function, see Function GetStackDisp (Getting Current Displacement Values) (p. 269).

2.14.1. Using Function RunCommand

This function enables you to issue a command from within a user subroutine. Inputs and outputs for RunCommand are as follows:

```
*deck,RunCommand
     function RunCommand (nChar, command)
c primary function:
                      Execute an ansys command
c object/library: upf
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
     nChar
            (int,sc,in)
                                    - Length of the command string (8 min)
C
      command (ch*(nChar),sc,in) - A character string containing a
С
                                              valid ANSYS command
C
c output arguments:
     RunCommand (int,sc,out)
                                    - An internally defined value, ignore
```

2.14.2. Using the /UNDO Command

The "undo" file you create by issuing the **/UNDO** command is similar to the File.DB file created when you issue the **SAVE** command. The **/UNDO** command format is:

```
/UNDO, Action
```

Action

ON, to write the undo file

OFF, to prevent the undo file from being written

PROMPT, to have the program ask permission before writing the file

STATUS, to restore the file as it existed after executing the last command issued before the **/UNDO** command.

2.14.3. Using the /HOLD command

Issue the /HOLD command to synchronize tasks. The program can synchronize tasks at the end of each results file set.

```
/HOLD,Filename,TimeInterval,Timeout
```

Filename

The eight-character name of a message file. If the named file exists, the program reads a command from the file and then deletes the file.

TimeInterval

The length of time, in seconds, that the program waits before trying to read the message file again.

Timeout

The maximum length of time, in seconds, that the program can wait between attempts to read the file.

Chapter 3: Accessing the Mechanical APDL Database

This chapter describes how you can retrieve information in the Mechanical APDL database (or store information in the database) by linking subroutines you create into the Mechanical APDL program.

You can use the database access routines with any of the user-programmable features. For example, you can create your own Mechanical APDL commands and use them to execute database access routines (or have a database access routine call a user-defined command).

Inputs and Outputs for Database Access Routines

The descriptions of the database access routines or functions within this chapter describe both the input arguments and output arguments. Argument information includes the argument's type, size and intent.

• Argument *type* is one of the following:

```
int - integer (4-byte)
long - integer (8-byte)
dp - double precision
log - logical
chr - character
comp - double precision complex
```

Argument size is one of the following:

```
sc - scalar variable ar(n) - array variable of length n func - functional return value
```

• Argument *intent* is one of the following:

```
in - input argumentout - output argumentinout - both an input and an output argument
```

Types of Database Access Routines

The rest of this chapter describes the functions and subroutines available for accessing information in the Mechanical APDL database. The function and subroutine descriptions are grouped into the following sections.

- 3.1. Routines for Selecting and Retrieving Nodes and Elements
- 3.2. Node Information Routines

- 3.3. Element Attribute Routines
- 3.4. Coupling and Constraint Routines
- 3.5. Nodal Loading Routines
- 3.6. Element Loading Routines
- 3.7. Results Information Routines

3.1. Routines for Selecting and Retrieving Nodes and Elements

3.1.1. Function ndnext (Getting the Next Node Number)

```
*deck,ndnext
     function ndnext (next)
c *** primary function: get the number of the next selected node
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
       next (int,sc,in) - the last node number used
С
                                   = 0 - use for initial value
С
    output arguments:
C
       ndnext (int,func,out) - the next selected node number
С
С
                                    = 0 - no more nodes
```

3.1.2. Function ndprev (Getting the Number of the Previous Selected Node)

```
*deck,ndprev
    function ndprev (next)
c *** primary function: get the number of the previous selected node
c *** Notice - This file contains ANSYS Confidential information ***
С
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
C
     input arguments:
С
     variable (typ,siz,intent) description
C
C
      next (int,sc,in)
                                - the next node number used
C
                                   = 0 - use for initial value
C
    output arguments:
С
       ndprev (int,func,out) - the previous selected node number
С
                                   = 0 - no more nodes
C
```

3.1.3. Function ndnxdf (Getting the Number of the Next Defined Node)

```
*deck,ndnxdf
    function ndnxdf (next)
c *** primary function: get the number of the next defined node
c
c *** Notice - This file contains ANSYS Confidential information ***
c
    input arguments:
```

```
c next (int,sc,in) - the last node number used
c = 0 - use for initial value
c
c output arguments:
c ndnxdf (int,func,out) - the next defined node number
c = 0 - no more nodes
```

3.1.4. Function ndsel (Selecting, Unselecting, Deleting, or Inverting a Node)

```
*deck,ndsel
     subroutine ndsel (ndmi,ksel)
c *** primary function: to select, unselect, delete, or invert a node.
c *** secondary functions: none.
c *** Notice - This file contains ANSYS Confidential information ***
      typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
     input arguments:
        variable (typ,siz,intent)
                                   description
С
                                   - node number
        ndmi (int,sc,in)
С
С
                                    = 0 - all nodes
С
                                     < 0 - do not delete CPs and CEQNs
                                           (merge/offset/compress)
C
                                   - type of operation to be performed.
С
        ksel
                (int,sc,in)
С
                                     ksel = 0 - delete node.
                                          = 1 - select node.
C
C
                                          =-1 - unselect node.
                                          = 2 - invert select status of node.
С
     output arguments:
С
C
        none.
```

3.1.5. Function elnext (Getting the Number of the Next Element)

```
*deck,elnext
     function elnext (next)
c *** primary function: get the number of the next selected element
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
       next (int,sc,in) - the last element number used
С
                                    = 0 - use for initial value
С
     output arguments:
      elnext (int,func,out)
                                 - the next selected element
С
С
                                    = 0 - no more elements
```

3.1.6. Function elprev (Getting the Number of the Previous Selected Element)

```
*deck,elprev
function elprev (prev)
c *** primary function: get the number of the previous selected element
c
```

```
c *** Notice - This file contains ANSYS Confidential information ***
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
С
С
    input arguments:
    variable (typ, siz, intent) description
C
      prev (int,sc,in) - the last element used
С
                                  = 0 - use for initial value
С
C
    output arguments:
     elprev (int,func,out) - the previous selected element
С
С
                                   = 0 - no more elements
```

3.1.7. Function elnxdf (Getting the Number of the Next Defined Element)

```
*deck,elnxdf
     function elnxdf (next)
c *** primary function: get the number of the next defined element
С
c *** Notice - This file contains ANSYS Confidential information ***
C
С
     input arguments:
      next (int,sc,in) - the last element number used
С
                                   = 0 - use for initial value
C
С
    output arguments:
С
      elnxdf (int,func,out) - the next defined element
С
                                   = 0 - no more elements
```

3.1.8. Subroutine elsel (Selecting, Unselecting, Deleting, or Inverting an Element)

```
*deck,elsel
    subroutine elsel (ielei,ksel)
c *** primary function: to select, unselect, delete, or invert an element.
c *** Notice - This file contains ANSYS Confidential information ***
```

3.2. Node Information Routines

3.2.1. Function ndingr (Getting Information About a Node)

The primary function of ndingr is getting information about a node. This function also sets the current node pointer to this node.

Note:

Some of the database commands in the input file shown below are in the common block ansysdef.inc, which must be included in the subroutine.

```
*deck,ndingr
      function ndingr (node, key)
c *** primary function: get information about a node.
c *** secondary functions: set current node pointer to this node.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
С
         node
                  (int,sc,in)
                                    - node number
                                       Should be 0 for key=11, DB_NUMDEFINED,
С
C
                                       DB_NUMSELECTED, DB_MAXDEFINED, and
С
                                       DB_MAXRECLENG
                                    - key as to information needed about
                  (int,sc,in)
C
         key
С
                                      the node.
С
                  = DB_SELECTED
                                   - return select status:
                      ndingr = 0 - node is undefined.
С
                              = -1 - node is unselected.
С
                              = 1 - node is selected.
С
                  = DB_NUMDEFINED - return number of defined nodes
C
                  = DB_NUMSELECTED - return number of selected nodes
C
                  = DB_MAXDEFINED - return highest node number defined
С
                  = DB_MAXRECLENG - return maximum record length (dp words)
С
                      2, return length (dp words)
С
                     3,
C
                     4, pointer to first data word
С
С
                  = 11, return void percent (integer)
С
                  = 17, pointer to start of index
С
                  = 117, return the maximum number of DP contact data stored for any node
С
                  = -1,
                  = -2, superelement flag
C
                  = -3, master dof bit pattern
С
C
                  = -4, active dof bit pattern
                     -5, solid model attachment
С
С
                  = -6, pack nodal line parametric value
                  = -7, constraint bit pattern
С
С
                  = -8, force bit pattern
С
                  = -9, body force bit pattern
                  = -10, internal node flag
С
С
                  = -11, orientation node flag =1 is =0 isnot
                  = -11, contact node flag <0
С
                  = -12, constraint bit pattern (for DSYM)
С
                  = -13, if dof constraint written to file.k (for LSDYNA only)
С
С
                  = -14, nodal coordinate system number (set by NROTATE)
С
                  =-101, pointer to node data record
С
                  =-102, pointer to angle record
                  =-103,
C
С
                  =-104, pointer to attached couplings
С
                  =-105, pointer to attacted constraint equations
                  =-106, pointer to nodal stresses
С
                  =-107, pointer to specified disp'S
С
                  =-108, pointer to specified forces
С
                  =-109, pointer to x/y/z record
C
                  =-110,
С
                  =-111.
C
C
                  =-112, pointer to nodal temperatures
                  =-113, pointer to nodal heat generations
C
С
                  =-115, pointer to calculated displacements
C
С
                  =-116,
С
      output arguments:
                (int,func,out) - the returned value of ndingr is based on
С
         ndingr
C
                                          setting of key.
```

3.2.2. Function getnod (Getting a Nodal Point)

```
*deck,getnod
     subroutine getnod (node, v, kerr, kcrot)
c *** primary function: get a nodal point
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
    node (int,sc,in)
                               - node number
С
С
     kerr (int,sc,inout) - message flag
                                 = 0 - print no message if node is unselected
C
С
                                       or undefined
                                 = 1 - print message if node is undefined
С
                                 = 2 - print message if node is undefined
С
C
                                       or unselected
                               - output coordinates in this coordinate system.
С
    kcrot (int,sc,in)
                                   if kcrot is negative, output theta and
С
                                   phi coordinates in radians
С
c output arguments:
           (dp,ar(6),out) - Coordinates (first 3 values) and rotation
C
                                   angles (last 3 values)
C
С
     kerr (int,sc,inout)
                               - select status
                                 = 0 - node is selected
С
                                  = 1 - node is not defined
C
                                  =-1 - node is unselected
```

3.2.3. Function putnod (Storing a Node)

```
*deck,putnod
     subroutine putnod (node, vctn, kcrot)
c *** primary function: store a node
c *** secondary functions: display node if in immediate mode.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
       node (int,sc,in) - node number
vctn (dp,ar(6),in) - array of 3 nodal coordinates and
С
                                              3 nodal rotation angles.
C
                                    - local coordinate system in which the nodal
С
        kcrot
                (int,sc,in)
С
                                       coordinates and angles are defined
     output arguments: none.
```

3.2.4. Function ndgall (Getting the XYZ/Rotation Coordinates Vector for a Node)

```
*deck,ndgall
    function ndgall (node,xyz)
c *** primary function:    get x,y,z,rotx,roty,rotz vector for a node.

c *** Notice - This file contains ANSYS Confidential information ***

c    input arguments:
        node    (int,sc,in) - node number for operation.
```

3.2.5. Subroutine ndspgt (Getting the Nodal Solution for a Node of an Element)

```
*deck,ndspqt
     subroutine ndspgt (node, dofs, ndof, nrot, xyzang, nuvect, unode)
c *** primary function: get the nodal solution for a node of an element
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
                                  - The node number
С
         node
                  (int,sc,in)
                  (int,ar(DOFBITLENG),in) - The dofs to retrieve for the node.
         dofs
C
                                     dof = degree of freedom
С
С
                                     The dofs array should be zeroed out,
C
                                     except for the needed parts.
                                     dofs is a bit pattern with true bits
С
                                     representing the GLOBAL Dof set desired.
С
                                     That is, dofs(1) is used for UX to SP06,
С
                                     and dofs(2) is used for TBOT to TTOP.
C
C
                                     See ECHPRM for details. For example,
                                           dofs(1) = UX + TEMP
С
                                           dofs(2) = TE3
С
                                     TTOP is a special case. If you want
С
С
                                     TTOP alone, use:
С
                                           dofs(2) = ibset(0,TTOP)
                                     If TBOT and TTOP are desired, you must use:
C
С
                                           dofs(2) = TBOT
                                           dofs(2) = ibset(dofs(2), TTOP)
С
         ndof
                  (int,sc,in)
                                  - The number of node dofs (1, 2 or 3).
С
         nrot
                  (int,sc,in)
                                  - Key to rotate dofs from nodal to global
С
                                     coordinate systems.
C
                                     if 0, none. if 2, 2-d. if 3, 3-d
С
С
                                     if > 0, dof set must include and only
                                     include all terms of the vector (e.g.
C
С
                                     UX, UY, UZ, or AX, AY, AZ).
                  (dp,ar(6),in) - The xyz virgin node coordinates
С
         xyzang
                                      (including angles). Not used if
С
                                      nrot = 0 or ndof < 2.
С
С
         nuvect.
                  (int,sc,in)
                                  - Number of vectors to retrieve. Can vary
                                     between 1 and 5. Normally 1 is what is
C
С
                                     wanted. Other vectors include previous
                                     values and/or velocities. See elucom for
C
С
                                     all possibilites. Contents are analysis
С
                                     type dependent.
      output arguments:
C
С
         unode
                  (dp,ar(ndof,nuvect),out) - Element nodal solution vectors in
                                               the global coordinate system.
C
```

3.3. Element Attribute Routines

3.3.1. Function elmigr (Getting Information About an Element)

```
*deck,elmiqr
     function elmiqr (ielem, key)
c *** primary function: get information about an element.
c *** secondary functions: set current element pointer to this element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
         ielem
                 (int,sc,in)
С
                                       should be zero for key=11, DB_NUMDEFINED,
С
                                         DB_NUMSELECTED, DB_MAXDEFINED, DB_MAXRECLENG,
C
                                    - information flag.
С
         key
                  (int,sc,in)
                                    - return select status:
С
                  = DB_SELECTED
                       elmiqr = 0 - element is undefined.
С
                               =-1 - element is unselected.
C
                               = 1 - element is selected.
С
                  = DB_NUMDEFINED - return number of defined elements
                                                                            (12)
С
                  = DB_NUMSELECTED - return number of selected elements
C
                                                                            (13)
С
                  = DB_MAXDEFINED - return maximum element number used
                                                                            (14)
                  = DB_MAXRECLENG - return maximum record length
C
                                                                            (15)
                                       (int words)
С
С
                  = 2 - return length (int words)
                  = 3 - return layer number
С
С
                        (for cross reference files return number of entities)
                  = 4 - return address of first data word
С
                  = 5 - return length (in record type units)
C
С
                  = 6 - return compressed record number.
С
                  = 11 - return void percent (integer)
                  = 16 - return location of next record
C
                         (this increments the next record count)
С
                  = 17 - pointer to start of index
С
                  = 18 - return type of file.
C
С
                      elmiqr = 0 - integer
                              = 1 - double precision
C
                              = 2 - real
= 3 - complex
С
С
                               = 4 - character*8
C
                               = 7 - index
С
С
                  = 19 - return virtual type of file.
С
                      elmiqr = 0 - fixed length (4.4 form)
                               = 1 - indexed variable length (layer data)
С
                               = 2 - xref data tables
С
                               = 3 - bitmap data (for 32 data item packed records)
С
                               = 4 - data tables (three dimensional arrays)
                  = 111 - return the maximum number of nodes stored for any element
C
С
                  = 123 - return the maximum number of DP contact data stored for any element
                  = -1 - material number
                                                   ( = -EL_MAT)
C
                    -2 - element type
                                                   ( = -EL_TYPE)
C
                    -3 - real constant number ( = -EL_REAL)
С
С
                    -4 - element section ID number ( = -EL_SECT)
                    -5 - coordinate system number ( = -EL_CSYS)
C
С
                           (see elmcmx for rest)
                  =-101 - pointer to element integers etc.
С
C
                              (see elmcmx with elmilg and 1 instead of -101)
C
      output arguments:
C
         elmiqr (int,sc,out) - the returned value of elmiqr is based on
C
                                        setting of key.
C
C
```

3.3.2. Function elmget (Getting an Element's Attributes and Nodes)

```
*deck,elmget
    function elmget (ielem,elmdat,nodes)
c *** primary function:    get element attributes and nodes.
c *** Notice - This file contains ANSYS Confidential information ***
```

3.3.3. Subroutine elmput (Storing an Element)

```
*deck,elmput
     subroutine elmput (ielem,elmdat,nnod,nodes)
c *** primary function: store element attributes and node numbers.
c *** secondary functions: set current element pointer to this element.
c *** Notice - This file contains ANSYS Confidential information ***
c *** NOTICE - The user is also responsible for defining the centroid for the
               element using the elmpct subroutine. Calling the elmput
C
С
               subroutine will NULL the element centroid previously defined.
     input arguments:
C
        ielem (int,sc,in)
                                    - element number
С
         elmdat.
                  (int,ar(EL_DIM),in) - element attributes.
С
                              elmdat(EL_MAT) - material number
    (EL_TYPE) - element type
C
С
                                     (EL REAL) - real constant number
С
                                     (EL_SECT) - section number
С
С
                                     (EL_CSYS) - coordinate system number
                                     (EL_DEAD) - death flag (bit 0)
C
С
                                            if clear - alive
                                            if set - dead
С
                                     (EL_SOLID) - solid model reference
C
                                     (EL_SHAPE) - 100*shape + specific shape
С
                                     (EL_OBJOPTIONS) - reserved
С
С
                                     (EL_PEXCLUDE) - p element include flag
С
                                             (bit 0)
                                             if clear - include
C
                                             if set - exclude
С
                                             For LSDYNA, it means part ID
С
                                            in regular ANSYS, it is never part ID
С
                                    - number of nodes for this element.
С
                  (int,sc,in)
                  (int,ar(*),in)
                                    - node numbers for this element.
С
         nodes
      output arguments: none.
```

3.3.4. Function etyiqr (Getting a Data Item About an Element Type)

```
*deck,etyiqr
function etyiqr (itype,key)
c *** primary function: get information about an element type.

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
c itype (int,sc,in) - element type number
c Should be 0 for key=11, DB_NUMDEFINED,
c DB_NUMSELECTED, DB_MAXDEFINED, and
c DB_MAXRECLENG
```

```
С
         key
                  (int,sc,in) - information flag.
                  = DB_SELECTED - return select status:
С
                       etyiqr = 0 - element type is undefined.
C
                              =-1 - element type is unselected.
C
С
                              = 1 - element type is selected.
                  = DB_NUMDEFINED - return number of defined element types
C
                  = DB_NUMSELECTED - return number of selected element types
С
                  = DB_MAXDEFINED - return highest element type number defined
С
                  = DB MAXRECLENG - return maximum record length (int words)
C
                 = -n, return element characteristic n from etycom for element
С
                       type itype.
С
                       n is correlated to the parameter names in echprm.
                       see elccmt for definitions of element characteristics.
                       note- this will not overwrite the current setting of
С
С
                        etycom.
С
     output arguments:
                                 - the returned value of etyiqr is based on
С
        etyiqr (int,func,out)
                                       setting of key.
С
```

3.3.5. Function etyget (Getting Information About an Element Type)

```
*deck,etyget
     function etyget (itype,ielx)
c *** primary function:
                         get element type data.
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
        itype (int,sc,in)
                                   - element type number
С
     output arguments:
С
С
        etyget (int,func,out)
                                   - status of element type.
С
                                     = 0 - element type is undefined.
                                     < 0 - number of data items on unselected
C
С
                                            element type.
                                     > 0 - number of data items on selected
С
                                            element type.
C
        ielx
                 (int,ar(*),out)
                                 - element type data. see elccmt for
                                     description of data.
c *** mpg etyget<el117,edgcntf1,edgcntsz,edgrde,edgrecc,edgmul:get elem type
```

3.3.6. Subroutine etyput (Storing Element Type Data)

```
*deck,etyput
     subroutine etyput (itype,n,ielx)
c *** primary function: store element type data.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
                                   - element type number for operation.
        itype (int,sc,in)
C
                                   - length of data vector to store.
С
                 (int,sc,in)
        ielx
С
                 (int,ar(*),in)
                                   - element type data. see elccmt for
C
                                     description.
     output arguments: none
```

3.3.7. Subroutine echrtr (Getting Information About Element Characteristics)

```
*deck,echrtr
     subroutine echrtr (iott,elcdn,ielc,kerr)
   primary function: collect all element characteristics based on
                                                    ityp, jtyp, and keyopts
c *** Notice - This file contains ANSYS Confidential information ***
C
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
С
c input arguments:
С
     variable (typ,siz,intent)
                                   description
                                     - printout file
С
     iott (int,sc,in)
             (int,ar(IELCSZ),inout) - input element characteristics
    ielc
C
                                     in positions 1 to 20.
С
С
                                     (itype, jstif, keyopts, etc.)
C
С
  output arguments:
    elcdn (chr,sc,out)
                                     - element descriptive name as character
С
                                       string
C
С
     ielc (int,ar(IELCSZ),inout) - input element characteristics
                                       in positions 21 to 150.
С
С
                                        (kdim, ishap, idegen, etc.)
                                        see elccmt for a full list
С
                                     - error flag
     kerr
              (int,sc,out)
С
                                       = 0 - no errors
С
                                       = 1 - errors
С
С
```

3.3.8. Subroutine etysel (Selecting, Unselecting, Deleting, or Inverting an Element Type)

```
*deck,etysel
     subroutine etysel (itypi,ksel)
c *** primary function: to select, unselect, delete, or invert an
                          element type.
c *** secondary functions: none.
C
c *** Notice - This file contains ANSYS Confidential information ***
С
C
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
     input arguments:
C
        variable (typ,siz,intent)
                                   description
С
С
        itypi (int,sc,in)
                                   - element type number
                                     = 0 - all element types
C
        ksel (int,sc,in)
                                   - type of operation to be performed.
С
С
                                     = 0 - delete element type.
C
                                     = 1 - select element type.
С
                                     =-1 - unselect element type.
                                     = 2 - invert element type.
С
C
С
     output arguments:
С
        none.
С
```

3.3.9. Function mpingr (Getting Information About a Material Property)

```
*deck,mpingr
              function mpingr (mat,iprop,key)
c *** primary function:
                                                                 get information about a material property.
c *** Notice - This file contains ANSYS Confidential information ***
               input arguments:
C
С
                      mat (int,sc,in)
                                                                                              - material number
С
                                                                                                     should be 0 for key=11,
                                                                                                      DB NUMDEFINED(12),
C
С
                                                                                                      DB_MAXDEFINED(14), and
                                                                                                      DB_MAXRECLENG(15)
С
                      iprop
                                           (int,sc,in)
                                                                                              - property reference number:
C
                        if iprop = 0, test for existence of any material property with this
С
                                                 material number (with key = DB_SELECTED(1))
С
                       ---- MP command labels -----
С
                      EX = 1, EY = 2, EZ = 3, NUXY = 4, NUYZ = 5, NUXZ = 6, GXY = 7, GYZ = 8
C
                       GXZ = 9, ALPX=10, ALPY=11, ALPZ=12, DENS=13, MU = 14, DAMP=15, KXX = 16
                       \texttt{KYY} = 17 \text{, } \texttt{KZZ} = 18 \text{, } \texttt{RSVX} = 19 \text{, } \texttt{RSVY} = 20 \text{, } \texttt{RSVZ} = 21 \text{, } \texttt{C} \qquad = 22 \text{, } \texttt{HF} \qquad = 23 \text{, } \texttt{VISC} = 24 \text
C
С
                       EMIS=25, ENTH=26, LSST=27, PRXY=28, PRYZ=29, PRXZ=30, MURX=31, MURY=32
                      MURZ=33, PERX=34, PERY=35, PERZ=36, MGXX=37, MGYY=38, MGZZ=39, EGXX=40
С
                      EGYY=41, EGZZ=42, SBKX=43, SBKY=44, SBKZ=45, SONC=46, DMPS=47, ELIM=48
C
                      USR1=49, USR2=50, USR3=51, USR4=51, FLUI=53, ORTH=54, CABL=55, RIGI=56
С
                      HGLS=57, BVIS=58, QRAT=59, REFT=60, CTEX=61, CTEY=62, CTEZ=63, THSX=64,
С
С
                       THSY=65, THSZ=66, DMPR=67, LSSM=68, BETD=69, ALPD=70, RH =71, DXX =72,
С
                       DYY =73, DZZ =74, BETX=75, BETY=76, BETZ=77, CSAT=78, CREF=79, CVH =80
С
                                          (see mpinit for uncommented code and for TB command information)
С
                       kev
                                              (int.sc.in)
                                                                                             - key as to the information needed
С
                                                                                                  about material property.
С
                                    = DB_SELECTED(1) - return select status:
С
                                                                mpinqr = 0 - material prop is undefined.
C
С
                                                                                  = 1 - material prop is selected.
                                    = DB_NUMDEFINED(12) - number of defined material properties
C
С
                                    = DB_MAXDEFINED(14) - highest material property number defined
                                    = DB_MAXRECLENG(15) - maximum record length (dp words)
                                   = 2 - return length (dp words)
C
                                    = 3 - return number of temp. values
С
                                    = 11 - return void percent (integer)
С
С
               output arguments:
                      mpinqr (int, func, out) - returned value of mpinqr is based on
С
                                                                                                     setting of key.
C
```

3.3.10. Function mpget (Getting a Material Property Table)

```
*deck,mpget
      function mpget (mat,iprop,temp,prop)
c *** primary function: get a material property table.
c *** Notice - This file contains ANSYS Confidential information ***
      typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                    intent=in,out,inout
C
С
      input arguments:
С
         variable (typ, siz, intent) description
                 (int,sc,in) - material number
(int,sc,in) - property reference number:
С
C
         iprop
         ---- MP command labels -----
C
```

```
EX = 1, EY = 2, EZ = 3, NUXY = 4, NUYZ = 5, NUXZ = 6, GXY = 7, GYZ = 8
C
        GXZ = 9, ALPX=10, ALPY=11, ALPZ=12, DENS=13, MU =14, DAMP=15, KXX =16
С
        KYY =17, KZZ =18, RSVX=19, RSVY=20, RSVZ=21, C =22, HF =23, VISC=24
C
        EMIS=25, ENTH=26, LSST=27, PRXY=28, PRYZ=29, PRXZ=30, MURX=31, MURY=32
С
С
        MURZ=33, PERX=34, PERY=35, PERZ=36, MGXX=37, MGYY=38, MGZZ=39, EGXX=40
        EGYY=41, EGZZ=42, SBKX=43, SBKY=44, SBKZ=45, SONC=46, DMPS=47, ELIM=48
C
         USR1=49, USR2=50, USR3=51, USR4=51, FLUI=53, ORTH=54, CABL=55, RIGI=56
С
        HGLS=57, BVIS=58, QRAT=59, REFT=60, CTEX=61, CTEY=62, CTEZ=63, THSX=64,
С
        THSY=65, THSZ=66, DMPR=67, LSSM=68, BETD=69, ALPD=70, RH =71, DXX =72,
C
        DYY =73, DZZ =74, BETX=75, BETY=76, BETZ=77, CSAT=78, CREF=79, CVH =80
С
С
                    (see mpinit for uncommented code and TB command information)
     output arguments:
С
                                   - number of temperature values
С
        mpget (int,func,out)
С
         temp
                 (dp,ar(mpget),out) - vector of the temperature values
                (dp,ar(mpget),out) - vector of the property values
        prop
```

3.3.11. Subroutine mpput (Storing a Material Property Table)

```
*deck, mpput
      subroutine mpput (mat,iprop,ntab,temp,prop)
c *** primary function:
                         store material property tables.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
C
        mat
                (int,sc,in)
                                    - material number.
                  (int,sc,in)
С
         iprop
                                    - property reference number:
         ---- MP command labels -----
C
         EX = 1, EY = 2, EZ = 3, NUXY = 4, NUYZ = 5, NUXZ = 6, GXY = 7, GYZ = 8
С
         GXZ = 9, ALPX=10, ALPY=11, ALPZ=12, DENS=13, MU =14, DAMP=15, KXX =16
C
         KYY =17, KZZ =18, RSVX=19, RSVY=20, RSVZ=21, C =22, HF =23, VISC=24
С
         {\tt EMIS=25,\ ENTH=26,\ LSST=27,\ PRXY=28,\ PRYZ=29,\ PRXZ=30,\ MURX=31,\ MURY=32}
С
         \texttt{MURZ=33, PERX=34, PERY=35, PERZ=36, MGXX=37, MGYY=38, MGZZ=39, EGXX=40}
С
С
         EGYY=41, EGZZ=42, SBKX=43, SBKY=44, SBKZ=45, SONC=46, DMPS=47, ELIM=48
         USR1=49, USR2=50, USR3=51, USR4=51, FLUI=53, ORTH=54, CABL=55, RIGI=56
С
        HGLS=57, BVIS=58, QRAT=59, REFT=60, CTEX=61, CTEY=62, CTEZ=63, THSX=64,
C
        THSY=65, THSZ=66, DMPR=67, LSSM=68, BETD=69, ALPD=70, RH =71, DXX =72,
С
        DYY =73, DZZ =74, BETX=75, BETY=76, BETZ=77, CSAT=78, CREF=79, CVH =80
С
С
С
                    (see mpinit for uncommented code and TB command information)
C
         ntab
                                    - number of entries in the table
                                       (1 to 100)
С
                 (dp,ar(ntab),in) - temperature vector (ascending)
C
         t.em
                 (dp,ar(ntab),in) - property vector
С
         prp
      output arguments:
C
         none.
```

3.3.12. Subroutine mpdel (Deleting a Material Property Table)

```
C
                                        iprop (int,sc,in)
                                                                                                                                                                - property reference number:
                                                                                                                                                                             (0 = all properties)
С
                                         ---- MP command labels -----
C
                                        EX = 1, EY = 2, EZ = 3, NUXY = 4, NUYZ = 5, NUXZ = 6, GXY = 7, GYZ = 8
С
С
                                        GXZ = 9, ALPX=10, ALPY=11, ALPZ=12, DENS=13, MU =14, DAMP=15, KXX =16
                                        \texttt{KYY} = 17 \text{, } \texttt{KZZ} = 18 \text{, } \texttt{RSVX} = 19 \text{, } \texttt{RSVY} = 20 \text{, } \texttt{RSVZ} = 21 \text{, } \texttt{C} \qquad = 22 \text{, } \texttt{HF} \qquad = 23 \text{, } \texttt{VISC} = 24 \text{, } \texttt{VISC} = 24 \text{, } \texttt{C} = 24 \text{, } \texttt{C}
C
                                          EMIS=25, ENTH=26, LSST=27, PRXY=28, PRYZ=29, PRXZ=30, MURX=31, MURY=32
С
                                        MURZ=33, PERX=34, PERY=35, PERZ=36, MGXX=37, MGYY=38, MGZZ=39, EGXX=40
С
                                        EGYY=41, EGZZ=42, SBKX=43, SBKY=44, SBKZ=45, SONC=46, DMPS=47, ELIM=48
C
                                        USR1=49, USR2=50, USR3=51, USR4=51, FLUI=53, ORTH=54, CABL=55, RIGI=56
                                        HGLS=57, BVIS=58, QRAT=59, REFT=60, CTEX=61, CTEY=62, CTEZ=63, THSX=64,
C
С
                                        THSY=65, THSZ=66, DMPR=67, LSSM=68, BETD=69, ALPD=70, RH =71, DXX =72,
  С
                                        DYY =73, DZZ =74, BETX=75, BETY=76, BETZ=77, CSAT=78, CREF=79, CVH =80
 C
 C
                                                                         (see mpinit for uncommented code and for TB command information)
  С
                           output arguments: none.
```

3.3.13. Function rlingr (Getting Information About a Real Constant Set)

```
*deck,rlingr
     function rlingr (nreal, key)
c *** primary function: get information about a real constant set
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
      typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                 intent=in.out.inout
  input arguments:
C
        variable (typ,siz,intent)
С
                                   description
        nreal (int,sc,in) - real constant table number
C
С
                                      should be 0 for key=11, DB_NUMDEFINED,
С
                                      DB_NUMSELECTED, DB_MAXDEFINED, and
                                      DB_MAXRECLENG
С
С
        key
                 (int,sc,in)
                                   - information flag.
С
                              - return number of values stored for nreal
             = DB_SELECTED - return select status
С
                      rlingr = 0 - real constant table is undefined.
С
                             =-1 - real constant table is unselected.
                             = 1 - real constant table is selected
С
             = DB_NUMDEFINED - return number of defined real constant tables
             = DB_NUMSELECTED - return number of selected real constant tables
С
C
             = DB_MAXDEFINED - return highest real constant table defined
             = DB_MAXRECLENG - return maximum record length (dp words)
c output arguments:
        rlingr (int,func,out) - the returned value of rlingr is based on
С
                                     setting of key.
C
c *** mpg magnetic interface usage
```

3.3.14. Function riget (Getting Real Constant Data)

```
*deck,rlget
     function rlget (nreal,rtable)
c *** primary function:    get real constant data

c *** Notice - This file contains ANSYS Confidential information ***

c    input arguments:
        nreal (int,sc,in) - real constant table number
```

```
c output arguments:
c    rlget (int,func,out) - number of real constant data obtained
c    rtable (dp,ar(*),out) - real constant data obtained
```

3.3.15. Subroutine rIsel (Selecting or Deleting a Real Constant Set)

```
*deck,rlsel
     subroutine rlsel (nreai, ksel)
c *** primary function: select or delete a real constant set
c *** secondary functions: none
С
c *** Notice - This file contains ANSYS Confidential information ***
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
C
 input arguments:
    variable (typ,siz,intent)
С
                               description
C
     nreai (int,sc,in)
                                - real constant table
                                  = 0 - all real constant tables
С
    ksel (int,sc,in)
                                - type of operation to be performed.
С
                                  = 0 - delete real constant table.
С
С
                                  = 1 - select real constant table.
С
                                  =-1 - unselect real constant table.
                                  = 2 - invert real constant table.
С
С
  output arguments:
C
С
     none
С
```

3.3.16. Function csyigr (Getting Information About a Coordinate System)

```
*deck,csyiqr
     function csyiqr (ncsy,key)
                         get information about a coordinate system
c *** primary function:
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
     ncsy (int,sc,in)
                                - coordinate system reference number
C
C
                                   should be zero for key= DB_NUMDEFINED
С
                                   or DB_MAXDEFINED
              (int,sc,in)
                                - information flag.
C
     key
               = DB_SELECTED - return status:
С
С
                            csyiqr = 0 - coordinate system is not defined
                                    -1 - coordinate system is not selected
C
                                     1 - coordinate system is selected
С
               = DB_NUMDEFINED - number of defined coordinate systems
С
С
               = DB_MAXDEFINED - maximum coordinate system reference
С
                                 number used.
С
  output arguments:
     csyiqr (int, func, out)
                                - the returned value of csyiqr is based on
С
                                   setting of key.
C
```

3.3.17. Function csyget (Getting a Coordinate System)

```
*deck,csyget
     function csyget (ncsy,csydpx,csyinx)
c *** primary function: get a coordinate system
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
  NOTE: As a time-saving device, this routine will not fetch the coordinate
С
          system data from the database (an expensive operation)
          if ncsy = csyinx(4), as this would indicate that the data is current.
C
С
          If you wish to force this routine to fetch coordinate system data (in
          the case of loading a local array, for example), you MUST set
С
         ncsy != csyinx(4) before function call.
C
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                   intent=in,out,inout
  input arguments:
С
C
     variable (typ,siz,intent) description
                                                               csycom name
     ncsy (int,sc,in)
                                - coordinate system number
C
     csyinx(4) (int,sc,inout) - coordinate system number
C
                                                               csyact
С
  output arguments:
С
     csydpx (dp,ar(18),out)
                          csydpx(1-9)
                                      - transformation matrix
C
                                (10-12) - origin (XC, YC, ZC)
С
                                (13-14) - coordinate system parameters cparm
С
С
                                                                       cparm2
С
                                (15)
                                       - spare
                                (16-18) - defining angles
С
     csyinx (int,ar(6),out)
С
                          csyinx(1-2) - theta, phi singularity keys
                                       - coordinate system type
                                                                       icdsvs
С
                                (3)
         (csyinx(4) is inout)
                                       - coordinate system number
С
                               (4)
                                                                       csyact
С
                                (5)
                                       - spare
                                       - spare
                                (6)
C
С
     csyget
              (int,func,out)
                                   - status of coordinate system
                                   = 0 - coordinate system exists
С
C
                                    = 1 - coordinate system doesn't exist
```

3.3.18. Subroutine csyput (Storing a Coordinate System)

```
*deck,csyput
      subroutine csyput (ncsy,csydpx,csyinx)
c *** primary function:
                          store a coordinate system
c *** Notice - This file contains ANSYS Confidential information ***
  input arguments:
C
      ncsy
              (int,sc,in)
                                 - coordinate system number
С
C
              (dp,ar(18),out)
С
                          csydpx(1-9)
                                         - transformation matrix
                                 (10-12) - origin (XC, YC, ZC)
С
                                 (13-14) - coordinate system parameters
C
                                                                            cparm
C
                                                                           cparm2
                                 (15)
С
                                        - spare
                                 (16-18) - defining angles
С
      csyinx
               (int,ar(6),out)
С
                          csyinx(1-2) - theta, phi singularity keys
С
С
                                (3) - coordinate system type
                                                                           icdsys
С
                                 (4)
                                      - coordinate system number
                                                                           csyact
                                      - spare
C
                                (5)
С
                                 (6)
                                     - spare
```

```
c output arguments: none
```

3.3.19. Subroutine csydel (Deleting a Coordinate System)

```
*deck,csydel
     subroutine csydel (ncsy)
c *** primary function:
                         delete a coordinate system
c *** secondary functions: none
C
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                intent=in.out.inout
C
c *** Notice - This file contains ANSYS Confidential information ***
С
c input arguments:
C
    variable (typ, siz, intent) description
    ncsy (int,sc,in) - coordinate system number
С
С
  output arguments:
С
C
    none
C
```

3.3.20. Subroutine userac (Demonstrates Use of Element Attribute Routines)

See Subroutine userac (Accessing Element Information) (p. 173) for an example that demonstrates how to use the userac subroutine to extract information about an element type and element real constants from the Mechanical APDL database. You can find this subroutine on your product-distribution media.

3.4. Coupling and Constraint Routines

3.4.1. Function cpinqr (Getting Information About a Coupled Set)

```
*deck.cpingr
     function cpinqr (ncp,key)
c *** primary function: get information about a coupled set
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
С
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
 input arguments:
C
    variable (typ,siz,intent) description
С
               (int,sc,in) - coupled set number
     ncp
C
                              - inquiry key:
С
     key
               (int,sc,in)
С
                                 should be zero for key=11, DB_NUMDEFINED,
                                  DB_NUMSELECTED, DB_MAXDEFINED, and
C
                                 DB_MAXRECLENG
                = DB_SELECTED - return select status
С
С
                       cpinqr = 1 - coupled set is selected
С
                                  = 0 - coupled set in undefined
                                  =-1 - coupled set in unselected
C
С
                = DB_NUMDEFINED - return number of defined coupled sets
                = DB_NUMSELECTED - return number of selected coupled sets
```

```
= DB_MAXDEFINED - return the number of the highest numbered
С
С
                                   coupled set
                = DB_MAXRECLENG - return length of largest coupled set record
С
                                  (max record length)
C
                               - return length (data units)
С
                = 2
                                - return layer number
                = 3
С
                = 4
                                - return address of first data word
С
                = 5
                                - return number of values stored for ncp
С
                = 11
                               - return void percent (integer)
C
                = 16
                               - return location of next record
                = -1
                               - return master node for this eqn (this is
С
С
                                 currently only used by solution DB object)
C
  output arguments:
С
    cpinqr (int,func,out) - the returned value of cpinqr is based on
С
С
                                      setting of key
С
```

3.4.2. Function cpget (Getting a Coupled Set)

```
*deck,cpget
     function cpget (ncp,ieqn)
c *** primary function: get a coupled set
c *** Notice - This file contains ANSYS Confidential information ***
 input arguments:
    ncp (int,sc,in)
                                          - coupled set number
C
С
c output arguments:
     cpget
                  (int,func,out)
                                        - number of nodes in list
С
                   (int,ar(cpget+2),out) - coupled set info:
С
                             ieqn(1:cpget) - list of coupled nodes
C
                             ieqn(cpget+1) - set degree of freedom
С
С
                             ieqn(cpget+2) - number of nodes in list
C
                                             (copy of return value)
```

3.4.3. Subroutine cpput (Storing a Coupled Set)

```
*deck,cpput
      subroutine cpput (ncp,n,ieqn)
c *** primary function: store a coupling set
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
    ncp (int,sc,in) - coupled set number
n (int,sc,in) - number of nodes in
               (int,sc,in) - number of nodes in coupled set
(int,ar(n+2),in) - info for storage
С
С
С
                              ieqn(1:n) - list of coupled nodes
                              ieqn(n+1) - degree of freedom label for set
С
С
    (ieqn(n+2) is inout) ieqn(n+2) - number of nodes in coupled set
                                            (copy of n)
c output arguments:
С
      ieqn(n+2) (int,sc,inout) - number of nodes in coupled set
C
                                         (another copy of n)
```

3.4.4. Subroutine cpsel (Selecting or Deleting a Coupled Set)

```
*deck,cpsel
     subroutine cpsel (ncpi,ksel)
c *** primary function: select or delete a coupled set
c *** secondary functions: none
c *** Notice - This file contains ANSYS Confidential information ***
С
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
С
c input arguments:
С
     variable (typ,siz,intent) description
               (int,sc,in)
     ncpi
                                - coupled set number
С
    ksel
              (int.sc.in)
                               - select/delete flag
C
                                 = 0 - delete coupled set
С
С
                                 = 1 - select coupled set
C
  output arguments:
С
     none
C
```

3.4.5. Function ceinqr (Getting Information About a Constraint Equation Set)

```
*deck,ceingr
     function ceinqr (nce, key)
c *** primary function: get information about a constraint equation set
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
                                   - constraint equation number
                 (int,sc,in)
     nce
C
С
     key
                 (int,sc,in)
                                   - inquiry key:
                                      should be zero for key=11, DB_NUMDEFINED,
C
С
                                       DB_NUMSELECTED, DB_MAXDEFINED, and
                                      DB_MAXRECLENG
            = DB SELECTED
                             - return select status
С
                                    ceinqr = 1 - equation is selected
C
С
                                           = 0 - equation is undefined
                                           =-1 - equation is unselected
C
             = DB_NUMDEFINED - return number of defined contraint equations
С
С
            = DB_NUMSELECTED - return number of selected contraint equations
            = DB_MAXDEFINED - return number of highest numbered constraint
C
                                equation defined
            = DB_MAXRECLENG - return length of longest contraint equation set
C
                                (max record length)
С
С
                             - return length (data units)
            = 3
                             - return layer number
C
                             - address of first data word
С
С
            = 5
                             - return number of values stored for nce
C
            = 11
                             - return void percent (integer)
                             - return location of next record
С
            = CE_NONLINEAR - return 1 if CE is nonlinear
С
            = CE_ELEMNUMBER - return associated element number
C
     output arguments:
С
        ceinqr (int,func,out)
                                   - the returned value of ceingr is based on
С
С
                                        setting of key
```

3.4.6. Function ceget (Getting a Constraint Equation)

```
*deck,ceget
function ceget (nce,ieqn,deqn)
c *** primary function: get a constraint equation
c *** Notice - This file contains ANSYS Confidential information ***
```

3.4.7. Subroutine ceput (Storing a Constraint Equation)

```
*deck,ceput
subroutine ceput (nce,n,ieqn,deqn)
c *** primary function: store a constraint equation
c *** Notice - This file contains ANSYS Confidential information ***
```

3.4.8. Subroutine cesel (Deleting or Selecting a Constraint Equation)

3.5. Nodal Loading Routines

The following table provides available body load keys for the routines used to access nodal body load information (ansNodeBodyLoadIqr, ansNodeBodyLoadGet, ansNodeBodyLoadPut, and ansNodeBodyLoadDel).

Table 3.1: Nodal Body Load Keys

Key	Internal Label	BF Cmd Label	Meaning
1	BODYLOAD_TEMP	TEMP	Temperature
2	BODYLOAD_FLUE	FLUE	Fluence
3	BODYLOAD_HGEN	HGEN	Heat generation
4	BODYLOAD_DGEN	DGEN	Diffusing substance genereation rate
5	BODYLOAD_MVDI	MVDI	Magnetic virtual displacement flags
6	BODYLOAD_CHRG	CHRDG	Magnetic virtual displacement flags
11	BODYLOAD_PORT	PORT	Interior waveguide/transmission line port

Key	Internal Label	BF Cmd Label	Meaning
14	BODYLOAD_VMEN	VMEN	Mean flow
16	BODYLOAD_IMPD	IMPD	Impedance sheet
18	BODYLOAD_FSOU	FSOU	Fluid flow source
20	BODYLOAD_VELO	VELO	Velocity
21	BODYLOAD_MASS	MASS	Mass source
22	BODYLOAD_SPRE	SPRE	Static pressure
23	BODYLOAD_FPBC	FPBC	Flouqet periodic phase
25	BODYLOAD_UFOR	UFOR	Force potential (same slot as EF)
26	BODYLOAD_HFLW	HFLW	Heat flow (same slot as EF)
27	BODYLOAD_SFOR	SFOR	Shear force (same slot as VELO)

3.5.1. Function disigr (Getting Information About Constraints)

```
*deck,disiqr
     function disigr (node, key)
c *** primary function: get information about constraints
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
С
        node
                (int,sc,in)
                                  - node number for inquire.
                 (int,sc,in)
                                 - key as to the information needed
        key
C
                           = 1
                                            - return constraint mask
С
                           = DB_MAXDEFINED,
С
                             DB_NUMDEFINED - return number of nodal constraints
C
С
                                               NOTE: both DB_MAXDEFINED and
                                               DB_NUMDEFINED produce the same
С
                                               functionality
С
      output arguments:
С
         disiqr (int,func,out)
                                 - the returned value of disigr is based on
С
С
                                     setting of key.
```

3.5.2. Function disget (Getting a Constraint at a Node)

```
*deck,disget
     function disget (inode,idf,value)
c *** primary function: get a constraint from the data base (in raw form)
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
     variable (typ,siz,intent) description
        inode
                (int,sc,in)
                                  - node number (negative value for no
C
                                                 partabeval)
С
        idf
                 (int,sc,in)
                                  - reference number for the DOF: (1-32)
С
   UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8
C
   AZ = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
С
    CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
C
    EMF = 25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
С
                                 (missing entries are spares)
```

```
С
      output arguments:
С
         disget (int,func,out)
                                   - status of constraint.
                                      = 0 - no constraint on this node
C
                                            for this DOF
C
С
                                      = 4 - this node has a constraint
                                             defined for this DOF
C
                                      = -4 - this node has a pseudo-support
С
С
                                             defined for this DOF
                                    - constraint values
        value
                (dp,ar(4),out)
C
                          value(1-2) - (real,imag) values of present settings
                          value(3-4) - (real,imag) values of previous settings
С
```

3.5.3. Subroutine disput (Storing a Constraint at a Node)

```
*deck,disput
     subroutine disput (node,idf,value)
c *** primary function:
                          store a constraint at a node.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
        node
                 (int,sc,in)
                                    - node number
С
                  (int,sc,in)
                                   - reference number of DOF: (1-32)
С
         UX = 1, UY = 2, UZ = 3, ROTX= 4, ROTY= 5, ROTZ= 6, AX = 7, AY = 8
C
         AZ = 9, VX = 10, VY = 11, VZ = 12, GFV1 = 13, GFV2 = 14, GFV3 = 15, WARP = 16
С
          CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
С
         EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
                (dp,ar(2),in)
                                  - (real, imag) values for constraint
С
        value
     output arguments: none.
```

3.5.4. Subroutine disdel (Deleting a Constraint at a Node)

```
*deck,disdel
     subroutine disdel (node,idf)
c *** primary function: delete a constraint at a node
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
        node (int,sc,in)
                                  - node number.
C
                                  - reference number of DOF: (1-32)
С
        idf
                 (int,sc,in)
         UX = 1, UY = 2, UZ = 3, ROTX= 4, ROTY= 5, ROTZ= 6, AX = 7, AY = 8
С
         AZ = 9, VX =10, VY =11, VZ =12, GFV1=13, GFV2=14, GFV3=15, WARP=16
С
         CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
С
         EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
     output arguments: none.
```

3.5.5. Function foriqr (Getting Information About Nodal Loads)

```
*deck,foriqr
function foriqr (node,key)
c *** primary function: get information about nodal loads.
```

```
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
С
        node
                (int,sc,in)
                                   - number of node being inquired about.
                                      should be 0 for key=DB_MAXDEFINED or
C
                                       DB_NUMDEFINED
С
С
        key
                  (dp,sc,in)
                                   - key as to information needed
                              = 1
C
                                          - return force mask for node
                              = DB_MAXDEFINED,
                               DB_NUMDEFINED - return number of nodal loadings
С
С
                                                  in model
С
                                    NOTE: both DB_MAXDEFINED and DB_NUMDEFINED
                                    produce the same functionality
C
С
     output arguments:
         foriqr (int,func,out)
                                   - the returned value of forigr is based on
С
С
                                      setting of key.
```

3.5.6. Function forget (Getting a Nodal Load at a Node)

```
*deck,forget
     function forget (inode,idf,value)
c *** primary function:
                         get a force from the data base (in raw form)
c *** Notice - This file contains ANSYS Confidential information ***
C
      input arguments:
        inode (int,sc,in)
                                   - node number (negative value for no
С
                                                  partabeval)
С
         idf
                  (int,sc,in)
                                   - reference number for the DOF: (1-32)
С
                                     (see echprm.inc)
С
С
     output arguments:
        forget (int,func,out) - status of constraint.
C
С
                                    = 0 - no loading on this node for this DOF
                                    = 4 - this node has a loading for this DOF
С
        value
                (dp,ar(4),out)
C
                          value(1-2) - (real,imag) values of present settings
С
С
                         value(3-4) - (real,imag) values of previous settings
```

3.5.7. Subroutine forput (Storing a Nodal Load at a Node)

```
*deck,forput
     subroutine forput (node,idf,value)
c *** primary function: store a nodal load at a node
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
С
        node
                (int,sc,in)
                                   - node number
        idf
                 (int,sc,in)
                                   - reference number for the DOF: (1-32)
         FX = 1, FY = 2, FZ = 3, MX = 4, MY = 5, MZ = 6, CSGX = 7, CSGY = 8
C
         CSGZ= 9, VFX =10, VFY =11, VFZ =12
С
         RATE=17
С
                           FLOW=19, HEAT=20, AMPS=21, FLUX=22, NPKE=23, NPDS=24
         CURT=25, VLTG=26
С
                                   (missing entries are spares)
                                   - (real, imag) values of force
C
        value
                (dp,ar(2),in)
    output arguments: none.
```

3.5.8. Subroutine fordel (Deleting a Nodal Load at a Node)

```
*deck,fordel
     subroutine fordel (node,idf)
c *** primary function: delete a nodal load at a node
c *** secondary functions: none.
c *** Notice - This file contains ANSYS Confidential information ***
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                 intent=in,out,inout
С
     input arguments:
С
     variable (typ,siz,intent) description
С
                (int,sc,in) - node number
(int,sc,in) - reference number for the DOF: (1-32)
С
       node
C
        FX = 1, FY = 2, FZ = 3, MX = 4, MY = 5, MZ = 6, CSGX= 7, CSGY= 8
        CSGZ= 9, VFX =10, VFY =11, VFZ =12
С
         RATE=17, FLOW=19, HEAT=20, AMPS=21, FLUX=22, NPKE=23, NPDS=24
С
        CURT=25, VLTG=26
                                   (missing entries are spares)
С
C
    output arguments:
       none.
```

3.5.9. Function ansNodeBodyLoadlqr (Getting Information About a Nodal Body Load)

```
*deck,ansNodeBodyLoadIqr
     function ansNodeBodyLoadIgr (BODYLOAD_KEY, node, key)
c *** primary function:
                        get information about a nodal body load
c *** Notice - This file contains ANSYS Confidential information ***
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func
                                                intent=in.out.inout
c input arguments:
c variable (typ, siz, intent) description
С
    BODYLOAD_KEY (int,sc,in) - body load key (ansysdef.inc)
    node (int,sc,in)
                                 - node number
C
                                    should be zero for key=2
C
                  (int,sc,in)
                                  - key for operation
С
                                    = 1 - return nodal body load status
С
                                    ansNodeBodyLoadIqr
C
С
                                     = 0 - node has no body load defined
                                     = 1 - node has a body load defined
С
С
                                     = 2 - return total number of nodal body
                                          load with BODYLOAD_KEY defined in model
c output arguments:
c ansNodeBodyLoadIqr (int,func,out) - the returned value of ansNodeBodyLoadIqr
                                  z is based on setting of key.
C
```

3.5.10. Function ansNodeBodyLoadGet (Getting a Nodal Body Load Value)

```
*deck,ansNodeBodyLoadGet function ansNodeBodyLoadGet (BODYLOAD_KEY,node,val)
```

```
c *** primary function: get specified nodal load
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
        BODYLOAD_KEY
                         (int,sc,in)
                                         - body load key (see ansysdef.inc)
C
        node
                           (int,sc,in)
                                         - node number
С
    output arguments:
C
       ansNodeBodyLoadGet (int,func,out) - nodal load status of node.
                                           = 0 - nodal load undefined
C
                                            = 1 - nodal load is defined
С
   val
                          (dp,ar(*),out) - the nodal load (new,old)
```

3.5.11. Subroutine ansNodeBodyLoadPut (Storing a Nodal Body Load)

```
*deck,ansNodeBodyLoadPut
      subroutine ansNodeBodyLoadPut (BODYLOAD_KEY,node,nval,val)
c *** primary function: store nodal body loads
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
         BODYLOAD_KEY(int,sc,in)
                                           - body load key (see ansysdef.inc)
C
                       (int,sc,in) - body load key (see ansystem)
(int,sc,in) - node number
(int,sc,in) - number of values to put
                (int,sc,in)
С
С
         nval
                      (dp ,ar(*),in) - nodal loads
         val
С
     output arguments: none.
```

3.5.12. Subroutine ansNodeBodyLoadDel (Deleting a Nodal Body Load)

```
*deck,ansNodeBodyLoadDel
     subroutine ansNodeBodyLoadDel (BODYLOAD_KEY, node)
c *** primary function: delete node body loads
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
С
C
     variable (typ, siz, intent) description
      BODYLOAD_KEY (int,sc,in) - body load key (see ansysdef.inc)
С
                     (int,sc,in)
                                      - node number
C
С
     output arguments:
С
       none
С
С
```

3.6. Element Loading Routines

The following table provides available body load keys for the routines used to access element body load information (ansElemBodyLoadIqr, ansElemBodyLoadGet, ansElemBodyLoadPut, and ansElemBodyLoadDel).

Table 3.2: Element Body Load Keys

Key	Internal Label	BFE Cmd Label	Meaning
1	BODYLOAD_TEMP	TEMP	Temperature
2	BODYLOAD_FLUE	FLUE	Fluence
3	BODYLOAD_HGEN	HGEN	Heat generation
4	BODYLOAD_DGEN	DGEN	Diffusing substance genereation rate
5	BODYLOAD_MVDI	MVDI	Magnetic virtual displacement flags
6	BODYLOAD_CHRG	CHRDG	Magnetic virtual displacement flags
7	BODYLOAD_JS	JS	Mass source/Current density
9	BODYLOAD_EF	EF	Electric field
12	BODYLOAD_FVIN	FVIN	Field volume interface flag
18	BODYLOAD_FSOU	FSOU	Fluid flow source
19	BODYLOAD_FORC	FORC	Body force density in momentum equation

The following table provides available surface load keys for the routines used to access element surface load information (ansElemSurfLoadIqr, ansElemSurfLoadGet, ansElemSurfLoadPut, and ansElemSurfLoadDel).

Table 3.3: Element Surface Load Keys

Key	Internal Label	Cmd Label	SF Cmd	SFE Cmd	Meaning				
LOADS									
1	SURFLOAD_PRES	PRES	х	х	Pressure				
2	SURFLOAD_CONV	CONV	х	х	Convection				
3	SURFLOAD_IMPD	IMPD	х	Х	Impedence				
4	SURFLOAD_DFLU	DFLU	Х	Х	Diffusing substance generation rate				
6	SURFLOAD_CHRG	CHRG	х	Х	Charge density				
7	SURFLOAD_PORT	PORT	х	х	Port				
8	SURFLOAD_RAD	RAD	х	Х	Radiation boundary				
9	SURFLOAD_FSIN	FSIN	х	х	FSI interface				
10	SURFLOAD_RDSF	RDSF	х	х	Surface-to-surface radiation				
11	SURFLOAD_SELV	SELV		х	Substructure				

Key	Internal Label	Cmd Label	SF Cmd	SFE Cmd	Meaning		
12	SURFLOAD_SHLD	SHLD	х	х	Surface normal velocity or acceleration		
13	SURFLOAD_FFLX	FFLX	х	х	Flow flux		
16	SURFLOAD_HFLU	HFLU	х	х	Heat flux (film coefficient in db)		
17	SURFLOAD_ATTN	ATTN	Х	х	Attenuation coefficient		
18	SURFLOAD_TIMP	TIMP	Х		Thermal impedance		
19	SURFLOAD_VIMP	VIMP	Х		Viscous impedance		
FLAGS							
1	SURFLOAD_MXWF	MXWF	х	х	Maxwell force		
2	SURFLOAD_FREE	FREE	Х	х	Free surface		
3	SURFLOAD_INF	INF	х	х	Infinite surface		
4	SURFLOAD_BLI	BLI	х		Boundary layer		
5	SURFLOAD_FSI	FSI	х	х	FSI interface		
9	SURFLOAD_RIGW	RIGW	х		Rigid wall		

3.6.1. Function ans ElemBodyLoadlqr (Getting Information About an Element Body Load)

```
*deck,ansElemBodyLoadIqr
      function ansElemBodyLoadIqr (BODYLOAD_KEY,ielem,key)
c *** primary function: get information about element loads.
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
c BODYLOAD_KEY (int,sc,in)
                              - body load key (see ansysdef.inc)
            (int,sc,in)
                              - element number
                                Should be 0 for key=11, DB_NUMDEFINED,
С
                                DB_MAXDEFINED, and DB_MAXRECLENG
C
c key
               (int,sc,in)
                              - information flag.
                = DB_SELECTED
                              - return status:
С
С
                                  ansElemBodyLoadIqr = 0 - element has no loads
С
                                                     = 1 - element has load defined
С
                = DB_NUMDEFINED - return number of loads defined for this element
С
                                  (rec length)
С
                = DB_MAXDEFINED - return number of loads defined in model
                = DB_MAXRECLENG - return maximum number of loads defined for
С
                                   any element (max rec length)
С
                = 2
С
                                 - return length (dp words)
                = 3
                                 - return layer number (for cross reference
C
                                  files returnnumber of entities)
С
                                 - return address of first data word
С
                = 4
С
                = 5
                                 - return length (dp words)
                = 6
                                 - return compressed record number.
С
                                 - return void percent (integer)
                =11
С
                                 - return location of next record (this
С
                =16
С
                                   increments the next record count)
                =18
                                  - return type of file.
С
                                  С
С
                                                     = 2 - real
C
С
                                                     = 3 - complex
                                                     = 4 - character*8
```

```
С
                                                        = 7 - index
                                   - return virtual type of file.
С
                =19
                                     ansElemBodyLoadIqr = 0 - fixed length (4.4 form)
C
                                                         = 1 - indexed variable length
C
С
                                                              (layer data)
                                                         = 2 - xref data tables
C
                                                         = 3 - bitmap data
С
                                                              (for 32 data item packed
С
C
                                                               records)
С
                                                         = 4 - data tables (three dimensional
C
                                                               arrays)
      output arguments:
С
         ansElemBodyLoadIqr (int,func,out)
                                               - the returned value of ansElemBodyLoadIqr is
С
С
                                                 based on setting of key.
```

3.6.2. Function ans ElemBodyLoadGet (Getting an Element Body Load Value)

```
*deck,ansElemBodyLoadGet
      function ansElemBodyLoadGet (BODYLOAD_KEY,ielem,val)
c *** primary function: get element loads (in raw form)
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
       BODYLOAD_KEY(int,sc,in) - body load key ielem (int,sc,in) - element number
                                        - body load key (see ansysdef.inc)
С
C
    output arguments:
С
       ansElemBodyLoadGet (int,func,out)
                                                - status of element.
C
                                                  = 0 - no element load
С
                                                  > 0 - number of element load
С
С
         val
                     (dp,ar(n,2),out) - the element load (new,old).
```

3.6.3. Subroutine ans ElemBodyLoadPut (Storing an Element Body Load)

```
*deck,ansElemBodyLoadPut
     subroutine ansElemBodyLoadPut (BODYLOAD_KEY,ielem,nVals,values)
c *** primary function: store element loads
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
       BODYLOAD_KEY(int,sc,in)
                                      - body load key (see ansysdef.inc)
C
        ielem (int,sc,in) - element number
nVals (int,sc,in) - number of element load values
С
       nVals
С
                    (dp,ar(nVals),in) - element load
С
С
    output arguments: none
```

3.6.4. Subroutine ans ElemBodyLoadDel (Deleting an Element Body Load)

```
*deck,ansElemBodyLoadDel subroutine ansElemBodyLoadDel (BODYLOAD_KEY,ielem)
```

3.6.5. Function ans Elem Surf Load Iqr (Getting Information About an Element Surface Load)

```
*deck,ansElemSurfLoadIqr
      function ansElemSurfLoadIqr (SURFLOAD_KEY,ielem,iface,key)
c *** primary function: get information about element pressure/convection
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
        SURFLOAD_KEY(int,sc,in)
С
                                       - surface load key
        ielem (int,sc,in)
C
                                       - element number
                                       should be zero for key=DB_NUMDEFINED or
С
                                       DB_MAXRECLENG
С
        iface
                    (int,sc,in)
                                       - face number for inquire (0-6)
C
                                       face number is needed for key=5. for
С
                                       other values of key, iface has different
С
С
                                       meaning (see below)
С
        key
                      (int,sc,in)
                                      - key as to the information needed
                      = 1
                                       - return pressure mask for element
C
С
                      = 5
                                       - return number of pressures for this
                                        element face. if face = 0,
С
C
                                         returns max. data size
С
                      = DB_NUMDEFINED,
                      = DB MAXDEFINED - return value is based on setting of iface
C
                                         NOTE: both DB_NUMDEFINED and
С
                                                DB_MAXDEFINED produce the same
С
C
                                                functionality
                                  iface = 0 - return number of surface loads defined
С
                                        = 1-6 - return number of pressure loads
С
                                                defined for this element.
C
С
                                                NOTE: only 1-6 is valid, but this
                                                       routine simply checks that iface
С
                                                       is in the range. The actual value
C
                                                       of iface does not matter in this case.
С
С
                     = DB_MAXRECLENG - return the maximum number of element
                                        pressures on any element (max record
C
                                        length)
С
С
      output arguments:
         ansElemSurfLoadIqr(int,func,out) - the returned value of ansElemSurfLoadIqr
С
C
                                             is based on setting of key.
С
```

3.6.6. Function ans Elem Surf Load Get (Getting an Element Surface Load Value)

```
*deck,ansElemSurfLoadGet
    function ansElemSurfLoadGet (SURFLOAD_KEY,elem,iface,value)
c *** primary function:    get an element face load
c *** Notice - This file contains ANSYS Confidential information ***
```

```
С
     input arguments:
       SURFLOAD_KEY
                           (int,sc,in)
                                              - surface load key
C
        elem
                            (int,sc,in)
                                              - element number (negative value for
C
С
                                                                no partabeval)
С
        iface
                             (int.sc.in)
                                              - face number (1-68)
С
     output arguments:
        ansElemSurfLoadGet (int,func,out)
                                              - status of element
C
С
                                                 =-1 - element has no surface load of given type
                                                 = 0 - element face has no surface load of given type
С
                                                 > 0 - number of surface load values defined
С
                                              - the element surface load values (real,imag)
С
        value
                             (dp ,ar(*),out)
                                                at the given face for the given type
```

3.6.7. Subroutine ans Elem Surf Load Put (Storing an Element Surface Load)

```
*deck,ansElemSurfLoadPut
      subroutine ansElemSurfLoadPut (SURFLOAD_KEY,ielem,iface,nval,
                                       value)
c *** primary function: store an element face loads.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
С
        SURFLOAD_KEY (int,sc,in)
                         (int,sc,in) - surface load key
(int,sc,in) - element number for operation
        ielem
C
                        (int,sc,in) - face number (1-68)
(int,sc,in) - number of values to
С
         nval
                                            - number of values to put
С
         value
                         (dp,ar(nval),in) - the element load (real,imag) at each
С
                                               face
С
C
     output arguments: none
```

3.6.8. Subroutine ans Elem Surf Load Del (Deleting an Element Surface Load)

```
*deck,ansElemSurfLoadDel
     subroutine ansElemSurfLoadDel (SURFLOAD_KEY,ielem,iface)
c *** primary function: delete a surface load on an element
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
        SURFLOAD_KEY (int,sc,in)
                                       - surface load key
С
        ielem
                       (int,sc,in)
                                       - element number
C
        iface
                      (int,sc,in)
                                       - face number
С
                                         = 0 - delete all pressures on this
С
С
С
                                         = 1-6 - delete pressure on this face
     output arguments: none.
С
С
```

3.7. Results Information Routines

3.7.1. Function dspiqr (Getting Information About Nodal Results)

```
*deck,dspiqr
     function dspiqr (node, key)
c *** primary function: get information about nodal results
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
        node
                (int,sc,in)
                                  - node number
C
                                     = 0 - return information based on the setting of key
С
                                     > 0 - return result mask for given node
С
        key
                (int,sc,in)
                                   - key as to the information needed
С
                                       when node > 0 and key = 1 --> return result mask for given node
                                       when node = 0 and key = DB_MAXRECLENG --> return maximum record length (dp
С
С
                                                                                 NOTE: only supported with databa
                                       when node = 0 and key = any other value --> return number of calculated di
С
С
     output arguments:
        dspiqr
                (int,func,out) - the returned value of dspiqr is based on setting of node/key
```

3.7.2. Function dspget (Getting a Nodal Result from the Database)

```
*deck,dspget
     function dspget (node,ndf,idf,value)
                            get a nodal result from the data base
c *** primary function:
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
         node (int,sc,in) - node number
C
         ndf
                   (int,sc,in)
                                       - number of results requested
         idf
                   (int,ary(ndf),in) - reference number for the DOF: (1-32)
С
     UX = 1, UY = 2, UZ = 3, ROTX = 4, ROTY = 5, ROTZ = 6, AX = 7, AY = 8 AZ = 9, VX = 10, VY = 11, VZ = 12, GFV1 = 13, GFV2 = 14, GFV3 = 15, WARP = 16
С
С
     CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
C
     EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
                                     (missing entries are spares)
С
     output arguments:
С
         value
                 (dp,ar(ndf),out) - result values
```

3.7.3. Subroutine dspput (Storing a Result at a Node)

```
c output arguments: none
```

3.7.4. Subroutine dspdel (Deleting a Result at a Node)

```
*deck,dspdel
      subroutine dspdel (node,ndf,idf)
c *** primary function:
                          delete a result at a node
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
        node (int,sc,in)
                                   - node number. (0 to delete DOF at all
C
С
                                     nodes)
        ndf
                 (int,sc,in)
                                   - number of DOFs to delete (0 to delete
С
                                     all DOFs)
C
        idf
                (int,ar(*),in)
                                   - reference number for the DOF: (1-32)
        UX = 1, UY = 2, UZ = 3, ROTX= 4, ROTY= 5, ROTZ= 6, AX = 7, AY = 8
С
С
         AZ = 9, VX = 10, VY = 11, VZ = 12, GFV1 = 13, GFV2 = 14, GFV3 = 15, WARP = 16
         CONC=17, HDSP=18, PRES=19, TEMP=20, VOLT=21, MAG =22, ENKE=23, ENDS=24
         EMF =25, CURR=26, SP01=27, SP02=28, SP03=29, SP04=30, SP05=31, SP06=32
С
     output arguments: none
```

3.7.5. Function emsiqr (Getting Information About an Element's Miscellaneous Summable Data)

```
*deck,emsiqr
     function emsigr (ielem,key)
c *** primary function:
                          get information about element misc summable data
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
        ielem
С
               (int,sc,in)
                                   - element number (or zero, see below)
С
         key
                 (int,sc,in)
                                   - key as to the information needed
                     = 1 - return info about misc summed data records
C
С
                                   ielem > 0 - return number of misc summed
С
                                                data items for this element
С
                                                 (record length)
                                          = 0 - return maximum number of misc
С
С
                                                 summed data items on any
                                                 element (max record length)
C
С
                      = DB_NUMDEFINED - return total number of misc summed data
                                        items defined in model
C
С
      output arguments:
        emsiqr (int,func,out)
                                   - the returned value of emsigr is based on
C
                                        setting of key
```

3.7.6. Function emsget (Getting an Element's Miscellaneous Summable Data)

```
*deck,emsget
function emsget (ielem,value)
c *** primary function: get element misc summable data.
```

```
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
С
        ielem (int,sc,in)

    element number

     output arguments:
С
       emsget (int,func,out) - status of element.
С
                                    = 0 - element is undefined
C
С
                                    > 0 - number of data items returned
                                  - element misc summed data.
      value (dp,ar(*),out)
C
                                  NOTE: the contents of this record is element
                                       dependent. See SMISC on ETABLE command
C
```

3.7.7. Subroutine emsput (Storing an Element's Miscellaneous Summable Data)

```
*deck,emsput
     subroutine emsput (ielem,nval,value)
c *** primary function: store misc. summable data for an element.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
       ielem (int,sc,in)
nval (int,sc,in)
                                    - element number
                                    - number of values to be stored
C
                (dp,ar(nval),in) - the misc summed data values
С
        value
    output arguements: none
C
С
                                    NOTE: the contents of this record is element
                                        dependent. See SMISC on ETABLE command
С
```

3.7.8. Subroutine emsdel (Deleting an Element's Miscellaneous Summable Data)

3.7.9. Function enfiqr (Getting Information About Element Nodal Forces)

```
*deck,enfiqr
    function enfiqr (ielem,key)
c *** primary function: get information about element nodal forces
```

```
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
        ielem
                 (int,sc,in)
                                    - element number (or zero, see below)
С
С
        key
                 (int,sc,in)
                                    - key as to the information needed
                     = 1 - return info about element nodal forces
C
                                    ielem > 0 - return number of element nodal
С
                                                  forces for this element
С
                                                  (record length)
C
                                          = 0 - return maximum number of element
                                                  nodal forces on any element
С
С
                                                  (max record length)
С
                      = DB_NUMDEFINED - return total number of element nodal
                                         forces defined in model
C
С
     output arguments:
         enfiqr (int,func,out)
                                    - the returned value of enfigr is based on
С
                                        setting of key
С
```

3.7.10. Function enfget (Getting an Element's Nodal Forces)

```
*deck,enfget
     function enfget (ielem, value)
c *** primary function:
                        get element nodal forces.
c *** Notice - This file contains ANSYS Confidential information ***
C
     input arguments:
                                   - element number
        ielem (int,sc,in)
С
     output arguments:
С
        enfget (int,func,out)
                                   - status of element.
C
С
                                     = 0 - element has no nodal forces
С
                                     > 0 - number of nodal forces returned
        value
               (dp,ar(*),out)
                                   - element nodal forces
C
```

3.7.11. Subroutine enfput (Storing an Element's Nodal Forces)

```
*deck,enfput
     subroutine enfput (ielem,nval,value)
c *** primary function: store nodal force results at an element.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
        ielem
                 (int,sc,in)
                                    - element number
С
                   (int,sc,in)
                                     - the total number of values
С
        nval
                                      NOTE: There may be a maximum of 3 sets of
С
С
                                     nodal forces in the record: static
                                     forces, inertia forces, and damping forces
C
С
        value
                 (dp,ar(nval),in) - nodal force results
     output arguments: none
C
```

3.7.12. Subroutine enfdel (Deleting an Element's Nodal Forces)

3.7.13. Function ensign (Getting Information About an Element's Nodal Stresses)

```
*deck,ensigr
     function ensigr (ielem, key)
c *** primary function: get information about element nodal stresses
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
                                   - element number (or zero, see below)
        ielem (int,sc,in)
C
С
         key
                 (int,sc,in)
                                   - key as to the information needed
                     = 1 - return info about element nodal stresses
С
                                    ielem > 0 - return number of element nodal
C
                                                  stresses for this element
С
                                                  (record length)
С
                                          = 0 - return maximum number of element
C
С
                                                  nodal stresses on any element
                                                  (max record length)
С
С
                      = DB_NUMDEFINED - return total number of element
                                        nodal stresses defined in model
C
     output arguments:
C
С
        ensigr (int,func,out)
                                   - the returned value of ensign is based on
                                        setting of key
С
```

3.7.14. Function ensget (Getting an Element's Nodal Stresses)

```
*deck,ensget
     function ensget (ielem, value)
c *** primary function: get element nodal stresses.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
        ielem (int,sc,in)
                                   - element number
C
     output arguments:
C
         ensget
                (int,func,out)
                                   - status of element.
С
                                     = 0 - element undefined
C
                                      > 0 - number of nodal stresses
С
С
                                           returned
C
        value
                 (dp,ar(*),out)
                                   - element nodal stresses
```

```
С
                                 NOTE: Stresses at each corner node in the order
С
                                          X, Y, Z, XY, YZ, XZ, S1, S2, S3, SI, SE
                                       For solid elements, stresses at each
C
С
                                          corner node
С
                                       For shell elements, stresses at each
                                          corner node (first top durface, then
C
                                          bottom)
С
                                       For layered elements (w/KEYOPT(8)=0),
С
                                          stresses for "first" layer at each
C
C
                                          corner node (first at the bottom
                                          surface of the bottom layer, then the
C
С
                                          top surface of the top layer).
C
                                          Stresses for "second" layer at each
                                          corner node (first the bottom surface,
C
                                          then the top surface for the layer with
С
С
                                          the largest failure criteria).
                                          The second layer is not present if
С
                                          failure criteria were not used or are
С
С
                                          not appropriate
                                       For layered elements (w/KEYOPT(8)=1),
С
                                          stresses for each layer at each corner
C
                                          node (first at the bottom surface, then
C
                                          the top surface)
С
                                       For beam elements, the contents of this
C
                                          record is element depenent. See LS
C
                                          item of ETABLE command.
C
```

3.7.15. Subroutine ensput (Storing Nodal Stresses at an Element)

```
*deck,ensput
      subroutine ensput (ielem,nval,value)
c *** primary function:
                           store nodal stresses at an element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
                                     - element number
С
        ielem
                (int,sc,in)
                                     - the total number of values
C
         nval
                   (int,sc,in)
                                       (11*nnod*nface)
С
С
         value
                (dp,ar(nval),in)
                                     - the stress values
С
      output arguments: none
C
                                NOTE: Stresses at each corner node in the order
                                         X, Y, Z, XY, YZ, XZ, S1, S2, S3, SI, SE
С
                                       For solid elements, stresses at each
C
С
С
                                       For shell elements, stresses at each
                                         corner node (first top surface, then
C
C
С
                                       For layered elements (w/KEYOPT(8)=0),
                                          stresses for "first" layer at each
C
С
                                          corner node (first at the bottom
                                          surface of the bottom layer, then the
C
С
                                          top surface of the top layer).
C
                                          Stresses for "second" layer at each
                                          corner node (first the bottom surface,
C
                                          then the top surface for the layer with
С
                                          the largest failure criteria).
С
                                         The second layer is not present if
С
                                          failure criteria were not used or are
С
C
                                         not appropriate
С
                                       For layered elements (w/KEYOPT(8)=1),
С
                                          stresses for each layer at each corner
                                          node (first at the bottom surface, then
C
                                          the top surface)
```

```
For beam elements, the contents of this record is element depenent. See LS item of ETABLE command.
```

3.7.16. Subroutine ensdel (Deleting an Element's Nodal Stresses)

3.7.17. Function esfiqr (Getting Information About Element Surface Stress Data)

```
*deck,esfigr
     function esfigr (ielem, key)
c *** primary function:
                         get information about element surface stress data
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
                                   - element number (or zero, see below)
        ielem (int,sc,in)
С
С
         key
                 (int,sc,in)
                                   - key as to the information needed
                 = 1 - return info about surface stress
С
                         ielem > 0 - return number of surface stresses on this
C
                                      element (rec length)
                                = 0 - return maximum number of surface stresses
С
                                      on any element (max rec length)
С
                  = DB_NUMDEFINED - return the number of surface stresses
С
                                    defined in model
C
     output arguments:
С
        esfigr (int,func,out) - the returned value of esfigr is based on
С
                                       setting of key
```

3.7.18. Function esfget (Getting Element Surface Stress Data)

```
*deck,esfget
    function esfget (ielem,value)
c *** primary function: get element surface stress data.

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
    ielem (int,sc,in) - element number

c output arguments:
```

3.7.19. Subroutine esfput (Storing Element Surface Stress Data)

```
*deck,esfput
      subroutine esfput (ielem,nval,value)
c *** primary function: store surface stresses for an element.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
       ielem (int,sc,in)
nval (int,sc,in)
                                     - element number
С
                                     - the total number of values
C
С
                                        (19 * number of stress faces)
                                        There is a max of 2 stress faces
С
                                     - the values
С
      value (dp,ar(nval),in)
     output arguments: none
```

3.7.20. Subroutine esfdel (Deleting an Element's Surface Stress Data)

3.7.21. Function engiqr (Getting Information About an Element's Energies)

```
*deck,engiqr
     function engiqr (ielem, key)
c *** primary function: get information about element energies
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
        ielem (int,sc,in)
                                   - element number (or zero, see below)
C
                                   - key as to the information needed
С
         key
                 (int,sc,in)
                 = 1 - return info about element energies
С
                              ielem > 0 - return number of element energies on
C
                                           this element (rec length)
С
С
                                    = 0 - return maximum number of element
                                           energies on any element
C
                                           (max rec length)
C
                  = {\tt DB\_NUMDEFINED} - return the number of element energies
С
С
                                     defined in model
```

3.7.22. Function engget (Getting an Element's Energies)

```
*deck, engget
      function engget (ielem, value)
c *** primary function:
                         get element energies.
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
        ielem (int,sc,in)
                                       - element number
C
      output arguments:
С
С
         engget (int,func,out)
                                       - status of element.
                                          = 0 - element undefined
С
                                          = MAXENG+1 - energies returned (see in echprm.inc)
C
                (dp,ar(MAXENG+1),out) - volume and energies
С
С
                                value(1) = volume of element
                                      (2) = stiffness energy
C
                                      (3) = artificial hourglass energy
С
С
                                      (4) = kinetic energy
                                      (5) = plastic energy
C
C
                                      (6) = creep energy
                                      (7) = stabilization energy
C
                                      (8) = strain energy density
С
                                      (9) = thermal energy
С
                                     (10) = viscous regularization energy for CZM
C
С
                                     (11) = sparse (future friction energy)
С
                                     (12) = damping energy
С
                                     (13) = external work by element load
С
                                     (14) = stiffness energy amplitude
                                     (15) = kinetic energy amplitude
С
                                     (16) = stiffness energy peak
C
                                     (17) = kinetic energy peak
С
С
                                     (18) = intermediate result for stif. energ. amplitude
                                            in harmonic analysis 1
С
С
                                     (19) = intermediate result for kin. energ. amplitude
                                            in harmonic analysis 1
C
С
                                     (20) = intermediate result for stif. energ. amplitude
С
                                            in harmonic analysis 2
                          (21 = MAXENG+1) = intermediate result for kin. energ. amplitude
C
                                            in harmonic analysis 2
```

3.7.23. Subroutine engput (Storing an Element's Energies and Volume)

```
*deck,engput
     subroutine engput (ielem,nval,value)
c *** primary function:
                          store volume and energies for an element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
С
        ielem (int,sc,in)
                                     - element number
              (int,sc,in)
                                     - the total number of values to be stored
С
        nval
                                       must be MAXENG (see in echprm.inc)
С
        value (dp,ar(MAXENG+1),in) - volume and energies
```

```
value(1) = volume of element
С
С
                                      (2) = stiffness energy
                                      (3) = artificial hourglass energy
C
                                      (4) = kinetic energy
С
С
                                      (5) = plastic energy
                                      (6) = creep energy
C
                                      (7) = stabilization energy
С
                                      (8) = strain energy density
С
                                      (9) = thermal energy
C
С
                                     (10) = viscous regularization energy for CZM
                                     (11) = sparse (future friction energy)
С
С
                                     (12) = damping energy
С
                                     (13) = external work by element load
                                     (14) = stiffness energy amplitude
C
                                     (15) = kinetic energy amplitude
С
С
                                     (16) = stiffness energy peak
                                     (17) = kinetic energy peak
C
                                     (18) = intermediate result for stif. energ. amplitude
С
С
                                            in harmonic analysis 1
                                     (19) = intermediate result for kin. energ. amplitude
C
                                            in harmonic analysis 1
C
                                     (20) = intermediate result for stif. energ. amplitude
C
                                            in harmonic analysis 2
С
                          (21 = MAXENG+1) = intermediate result for kin. energ. amplitude
С
                                            in harmonic analysis 2
C
С
      output arguments: none
```

3.7.24. Subroutine engdel (Deleting an Element's Energies)

3.7.25. Function egriqr (Getting Information About an Element's Nodal Gradients)

```
*deck,egrigr
      function egriqr (ielem, key)
c *** primary function: get information about element nodal gradients
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
                                   - element number (or zero, see below)
С
        ielem (int,sc,in)
                                   - key as to the information needed
С
                 (int,sc,in)
С
                 = 1 - return info about nodal gradients
                          for ielem > 0 - return number of nodal gradients on
С
С
                                            this element (record length)
С
                                    = 0 - return maximum number of nodal
                                            gradients on any element
C
                                            (maximum record length)
```

3.7.26. Function egrget (Getting an Element's Nodal Gradients)

```
*deck,egrget
     function egrget (ielem, value)
c *** primary function:
                         get element nodal gradients.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
С
        ielem (int,sc,in)
                                   - element number
С
     output arguments:
        egrget (int,func,out)
                                    - status of element.
С
                                     = 0 - element undefined
С
                                      > 0 - number of nodal gradients
С
С
                                             returned
       value
                (dp,ar(*),out)
                                    - element nodal gradients
C
С
                                     Note: If a coupled field, a set of
                                       gradients are stored in the following
C
С
                                       order (as available): fluid, thermal,
                                       electric, magnetic
C
c *** mpg egrget < pagend, magget < hsnget2: get elem gradient, H,
```

3.7.27. Subroutine egrput (Storing an Element's Nodal Gradients)

```
*deck,egrput
     subroutine egrput (ielem,nval,value)
c *** primary function: store nodal gradients at an element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
        ielem (int,sc,in)
                                    - element number
C
        nval
                  (int,sc,in)
                                    - the total number of values
С
                                      (ndir*nnod*nscalr)
С
        value (dp,ar(nval),in)
                                    - the gradient values
C
                                    Note: If a coupled field, a set of
С
                                      gradients are stored in the following
С
С
                                      order (as appropriate): fluid, thermal,
                                      electric, magnetic
C
     output arguments: none
```

3.7.28. Subroutine egrdel (Deleting an Element's Nodal Gradients)

```
*deck,egrdel
```

3.7.29. Function eeliqr (Getting Information About an Element's Nodal Elastic Strains)

```
*deck,eeliqr
      function eeligr (ielem, key)
c *** primary function: get information about element nodal elastic strains
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
         ielem (int,sc,in) - element number (or zero, see below)
key (int,sc,in) - key as to the information needed
C
С
                  = 1 - return info about elastic strains
С
                                ielem > 0 - return number of nodal elasic strains
C
                                              on this element (rec length)
                                      = 0 - return maximum number of nodal elastic
С
С
                                              strains on any element
С
                                              (max rec length)
                   = DB_NUMDEFINED - return the number of nodal elastic strains
C
С
                             defined in model
      output arguments:
C
С
         eeliqr (int,func,out)
                                      - the returned value of eeligr is based on
                                           setting of key
С
```

3.7.30. Function eelget (Getting an Element's Nodal Elastic Strains)

```
*deck,eelget
     function eelget (ielem, value)
                         get element nodal elastic strains.
c *** primary function:
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
        ielem (int,sc,in)
C
                                   - element number
С
     output arguments:
                                   - status of element.
        eelget (int,func,out)
C
C
                                      = 0 - element undefined
                                      > 0 - number of nodal elastic strains
С
                                            returned
C
        value
                  (dp,ar(*),out)
                                   - element nodal elastic strains
С
С
                                NOTE: Strains at each corner node in the order
                                        X, Y, Z, XY, YZ, XZ, EQV
C
С
                                      For solid elements, strains at each
С
                                         corner node
                                      For shell elements, strains at each
C
                                        corner node (first top durface, then
```

```
C
                                          bottom)
С
                                       For layered elements (w/KEYOPT(8)=0),
                                          strains for "first" layer at each
C
                                          corner node (first at the bottom
С
С
                                          surface of the bottom layer, then the
                                          top surface of the top layer).
C
                                          Strains for "second" layer at each
С
                                          corner node (first the bottom surface,
С
                                          then the top surface for the layer with
C
C
                                          the largest failure criteria).
                                          The second layer is not present if
C
С
                                          failure criteria were not used or are
C
                                          not appropriate
                                       For layered elements (w/KEYOPT(8)=1),
C
                                          strains for each layer at each corner
С
С
                                          node (first at the bottom surface, then
С
                                          the top surface)
                                       For beam elements, the contents of this
С
С
                                          record is element depenent. See LEPEL
                                          item of ETABLE command.
С
```

3.7.31. Subroutine eelput (Storing an Element's Nodal Elastic Strains)

```
*deck,eelput
     subroutine eelput (ielem,nval,value)
c *** primary function:
                           store nodal elastic strains at an element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
                                      - element number
        ielem
                 (int,sc,in)
C
С
         nval
                   (int,sc,in)
                                      - the total number of values
                                       (7*nnod*nface)
С
         value
                  (dp,ar(nval),in)
                                     - nval strain values
C
      output arguments: none
С
                                NOTE: Strains at each corner node in the order
C
С
                                          X, Y, Z, XY, YZ, XZ, EQV
                                       For solid elements, strains at each
C
С
                                          corner node
                                       For shell elements, strains at each
C
С
                                          corner node (first top durface, then
С
                                         bottom)
                                       For layered elements (w/KEYOPT(8)=0),
C
                                          strains for "first" layer at each
С
                                          corner node (first at the bottom
С
                                          surface of the bottom layer, then the
C
                                          top surface of the top layer).
С
                                          Strains for "second" layer at each
С
                                          corner node (first the bottom surface,
C
С
                                          then the top surface for the layer with
                                          the largest failure criteria).
C
С
                                         The second layer is not present if
C
                                          failure criteria were not used or are
                                         not appropriate
C
                                       For layered elements (w/KEYOPT(8)=1),
С
С
                                          strains for each layer at each corner
С
                                          node (first at the bottom surface, then
                                          the top surface)
С
                                       For beam elements, the contents of this
C
С
                                          record is element depenent. See LEPEL
С
                                          item of ETABLE command.
```

3.7.32. Subroutine eeldel (Deleting an Element's Nodal Elastic Strains)

3.7.33. Function epliqr (Getting Information About an Element's Nodal Plastic Strains)

```
*deck,epliqr
     function epliqr (ielem, key)
c *** primary function: get information about element nodal plastic strains
c *** Notice - This file contains ANSYS Confidential information ***
С
      input arguments:
                                  element number (or zero, see below)key as to the information needed
        ielem (int,sc,in)
C
         key
                 (int,sc,in)
C
                  = 1 - return info about plastic strains
С
                              ielem > 0 - return number of nodal plastic strains
C
                                             on this element
С
                                             (record length)
С
                                     = 0 - return maximum number of nodal plastic
C
С
                                             strains on any element
                                             (max record length)
С
С
                  = DB_NUMDEFINED - return the number of nodal plastic strains
                                      defined in model
C
      output arguments:
C
С
         epliqr (int,func,out) - the returned value of epliqr is based on
                                         setting of key
C
```

3.7.34. Function eplget (Getting an Element's Nodal Plastic Strains)

```
*deck,eplget
     function eplget (ielem, value)
c *** primary function: get element nodal plastic strains.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
       ielem (int,sc,in)
                                   - element number
С
     output arguments:
С
                (int,func,out)
                                   - status of element.
С
        eplget
                                     = 0 - element undefined
С
                                     > 0 - number of nodal plastic strains
С
С
                                           returned
C
        value
               (dp,ar(*),out)
                                   - element nodal plastic strains
```

```
C
                                 NOTE: Strains at each corner node in the order
                                          X, Y, Z, XY, YZ, XZ, EQV
С
                                       For solid elements, strains at each
C
С
                                          corner node
С
                                       For shell elements, strains at each
                                          corner node (first top durface, then
C
                                          bottom)
С
                                       For layered elements (w/KEYOPT(8)=0),
С
                                          strains for "first" layer at each
C
                                          corner node (first at the bottom
                                          surface of the bottom layer, then the
C
С
                                          top surface of the top layer).
                                          Strains for "second" layer at each
C
                                          corner node (first the bottom surface,
C
                                          then the top surface for the layer with
С
                                          the largest failure criteria).
С
                                          The second layer is not present if
С
                                          failure criteria were not used or are
С
С
                                          not appropriate
                                       For layered elements (w/KEYOPT(8)=1),
С
                                          strains for each layer at each corner
                                          node (first at the bottom surface, then
C
                                          the top surface)
С
                                       For beam elements, the contents of this
                                          record is element depenent. See LEPPL
C
                                          item of ETABLE command.
C
```

3.7.35. Subroutine eplput (Storing an Element's Nodal Plastic Strains)

```
*deck,eplput
      subroutine eplput (ielem,nval,value)
c *** primary function:
                          store nodal plastic strains at a element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
                                     - element number
С
        ielem
                (int,sc,in)
        nval
                                      - the total number of values
C
                   (int.sc.in)
                                       (6*nnod*nface)
С
С
        value
                (dp,ar(nval),in)
                                     - the strain values
С
      output arguments: none
                                NOTE: Strains at each corner node in the order
C
С
                                         X, Y, Z, XY, YZ, XZ, EQV
                                      For solid elements, strains at each
С
                                          corner node
C
                                       For shell elements, strains at each
С
С
                                          corner node (first top durface, then
C
                                          bottom)
                                      For layered elements (w/KEYOPT(8)=0),
C
С
                                          strains for "first" layer at each
                                          corner node (first at the bottom
C
                                          surface of the bottom layer, then the
C
                                          top surface of the top layer).
C
                                          Strains for "second" layer at each
С
C
                                          corner node (first the bottom surface,
                                          then the top surface for the layer with
C
                                          the largest failure criteria).
С
                                         The second layer is not present if
С
С
                                          failure criteria were not used or are
С
                                         not appropriate
                                      For layered elements (w/KEYOPT(8)=1),
C
С
                                          strains for each layer at each corner
С
                                          node (first at the bottom surface, then
C
                                          the top surface)
                                      For beam elements, the contents of this
```

```
c record is element depenent. See LEPPL item of ETABLE command.
```

3.7.36. Subroutine epidel (Deleting an Element's Nodal Plastic Strains)

3.7.37. Function ecriqr (Getting Information About an Element's Nodal Creep Strains)

```
*deck,ecriqr
     function ecriqr (ielem, key)
c *** primary function: get information about element nodal creep strains
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
        ielem (int,sc,in) - element number (or zero, see below)
key (int,sc,in) - key as to the information needed
С
         key (int,sc,in)
С
С
                  = 1 - return info about creep strains
                               ielem > 0 - return number of nodal creep strains
С
                                              on this element
C
                                              (record length)
С
                                      = 0 - return maximum number of nodal creep
С
С
                                              strains on any element
С
                                              (max record length)
                   = DB_NUMDEFINED - return the number of nodal creep strains
C
С
                                      defined in model
     output arguments:
С
         ecriqr (int, func, out) - the returned value of ecriqr is based on
С
С
                                          setting of key
```

3.7.38. Function ecrget (Getting an Element's Nodal Creep Strains)

```
*deck,ecrget
    function ecrget (ielem,value)
c *** primary function: get element nodal creep strains.

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
c ielem (int,sc,in) - element number

c output arguments:
c ecrget (int,func,out) - status of element.
```

```
С
                                       = 0 - element undefined
С
                                       > 0 - number of nodal creep strains
                                              returned
C
С
         value
                  (dp,ar(*),out)
                                     - element nodal creep strains
                                 NOTE: Strains at each corner node in the order
C
                                          X, Y, Z, XY, YZ, XZ, EQV
С
                                       For solid elements, strains at each
С
                                          corner node
C
С
                                       For shell elements, strains at each
                                          corner node (first top durface, then
C
С
C
                                       For layered elements (w/KEYOPT(8)=0),
                                          strains for "first" layer at each
C
                                          corner node (first at the bottom
С
С
                                          surface of the bottom layer, then the
                                          top surface of the top layer).
С
                                          Strains for "second" layer at each
С
С
                                          corner node (first the bottom surface,
                                          then the top surface for the layer with
С
                                          the largest failure criteria).
C
                                          The second layer is not present if
C
                                          failure criteria were not used or are
С
                                          not appropriate
C
                                       For layered elements (w/KEYOPT(8)=1),
C
                                          strains for each layer at each corner
С
С
                                          node (first at the bottom surface, then
C
                                          the top surface)
                                       For beam elements, the contents of this
С
                                          record is element depenent. See LEPCR
С
                                          item of ETABLE command.
C
```

3.7.39. Subroutine ecrput (Storing an Element's Nodal Creep Strains)

```
*deck,ecrput
      subroutine ecrput (ielem,nval,value)
c *** primary function:
                           store nodal creep strains at an element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
С
         ielem
                   (int,sc,in)
                                      - element number
                                      - the total number of values
         nval
                   (int,sc,in)
C
C
                                        (6*nnod*nface)
         value
                  (dp,ar(nval),in)
                                     - the strain values
С
      output arguments: none
С
С
                                NOTE: Strains at each corner node in the order
                                         X, Y, Z, XY, YZ, XZ, EQV
C
                                       For solid elements, strains at each
C
С
                                          corner node
                                       For shell elements, strains at each
C
С
                                          corner node (first top durface, then
                                          bottom)
C
С
                                       For layered elements (w/KEYOPT(8)=0),
C
                                          strains for "first" layer at each
                                          corner node (first at the bottom
C
                                          surface of the bottom layer, then the
С
                                          top surface of the top layer).
С
С
                                          Strains for "second" layer at each
                                          corner node (first the bottom surface,
С
                                          then the top surface for the layer with
C
С
                                          the largest failure criteria).
С
                                          The second layer is not present if
                                          failure criteria were not used or are
C
                                          not appropriate
С
```

```
For layered elements (w/KEYOPT(8)=1),
strains for each layer at each corner
node (first at the bottom surface, then
the top surface)

For beam elements, the contents of this
record is element depenent. See LEPCR
item of ETABLE command.
```

3.7.40. Subroutine ecrdel (Deleting an Element's Nodal Creep Strains)

3.7.41. Function ethiqr (Getting Information About an Element's Nodal Thermal Strains)

```
*deck,ethigr
     function ethiqr (ielem, key)
c *** primary function: get information about element nodal thermal strains
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
                                - element number (or zero, see below)
       ielem (int,sc,in)
        key
                                   - key as to the information needed
C
                 (int,sc,in)
                 = 1 - return info about thermal strains
С
С
                             ielem > 0 - return number of nodal thermal strains
                                           on this element.
C
С
                                           (record length)
                                   = 0 - return maximum number of nodal thermal
С
C
                                           strains on any element
                                           (max record length)
С
                  = DB_NUMDEFINED - return the number of nodal thermal strains
С
                                    defined in model
C
     output arguments:
C
С
        ethigr (int,sc,out)
                                   - the returned value of ethigr is based on
С
                                       setting of key
```

3.7.42. Function ethget (Getting an Element's Nodal Thermal Strains)

```
*deck,ethget
function ethget (ielem,value)

c *** primary function: get element nodal thermal strains.

c also the volumetric swelling strain

c *** Notice - This file contains ANSYS Confidential information ***
```

```
С
      input arguments:
         ielem
                 (int,sc,in)
                                     - element number
С
      output arguments:
С
С
         ethget (int,func,out)
                                     - status of element.
                                       = 0 - element undefined
С
                                       > 0 - number of nodal thermal strains
С
С
                                              returned
         value
                                     - element nodal thermal strains
C
                  (dp,ar(*),out)
                                 NOTE: Strains at each corner node in the order
C
С
                                          X, Y, Z, XY, YZ, XZ, EQV, epswel
C
                                       For solid elements, strains at each
                                          corner node
C
                                       For shell elements, strains at each
С
С
                                          corner node (first top durface, then
С
                                          bottom)
                                       For layered elements (w/KEYOPT(8)=0),
С
                                          strains for "first" layer at each
С
                                          corner node (first at the bottom
С
                                          surface of the bottom layer, then the
C
                                          top surface of the top layer).
C
                                          Strains for "second" layer at each
С
                                          corner node (first the bottom surface,
С
                                          then the top surface for the layer with
C
                                          the largest failure criteria).
С
С
                                          The second layer is not present if
                                          failure criteria were not used or are
C
                                          not appropriate
С
                                       For layered elements (w/KEYOPT(8)=1),
С
                                          strains for each layer at each corner
С
                                          node (first at the bottom surface, then
C
C
                                          the top surface)
                                       For beam elements, the contents of this
С
С
                                          record is element depenent. See LEPTH
                                          item of ETABLE command.
C
```

3.7.43. Subroutine ethput (Storing an Element's Nodal Thermal Strains)

```
*deck.et.hput
      subroutine ethput (ielem,nval,value)
c *** primary function:
                          store nodal thermal strains at an element.
                            also the volumetric swelling strain
C
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
С
        ielem
                (int,sc,in)
                                     - element number
                                      - the total number of values
C
         nval
                   (int,sc,in)
                                       (7*nnod*nface)
С
С
         value
                  (dp,ar(nval),in)
                                     - the strain values
      output arguments: none
С
                                 NOTE: Strains at each corner node in the order
C
                                         X, Y, Z, XY, YZ, XZ, EQV, epswel
С
                                       For solid elements, strains at each
С
                                          corner node
C
                                       For shell elements, strains at each
С
                                          corner node (first top durface, then
С
С
                                          bottom)
                                       For layered elements (w/KEYOPT(8)=0),
С
                                          strains for "first" layer at each
C
С
                                          corner node (first at the bottom
С
                                          surface of the bottom layer, then the
                                          top surface of the top layer).
C
                                          Strains for "second" layer at each
С
```

```
С
                                          corner node (first the bottom surface,
С
                                          then the top surface for the layer with
                                          the largest failure criteria).
C
                                         The second layer is not present if
C
С
                                          failure criteria were not used or are
                                         not appropriate
C
                                       For layered elements (w/KEYOPT(8)=1),
С
                                         strains for each layer at each corner
С
                                         node (first at the bottom surface, then
C
С
                                         the top surface)
                                       For beam elements, the contents of this
С
С
                                         record is element depenent. See LEPTH
                                          item of ETABLE command.
```

3.7.44. Subroutine ethdel (Deleting an Element's Thermal, Initial, and Swelling Strains)

3.7.45. Function euligr (Getting Information About an Element's Euler Angles)

```
*deck.euligr
     function euligr (ielem, key)
c *** primary function: get information about element euler angles
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
         ielem (int,sc,in)
                                    - element number (or zero, see below)
С
                                - element number (02 - ) - key as to the information needed
С
         key
                   (int,sc,in)
                  = 1 - return info about element euler angles
C
                              ielem > 0 - return number of euler angles on this
С
                                             element.
С
                                             (record length)
С
                                     = 0 - return maximum number of euler angles
С
                                             on any element
C
C
                                             (max record length)
                  = DB_NUMDEFINED - return the number of element euler angles
С
                                     defined in model
С
С
      output arguments:
         euligr (int,func,out) - the returned value of euligr is based on
С
                                         setting of key
С
```

3.7.46. Function eulget (Getting an Element's Nodal Euler Angles)

```
*deck,eulget
     function eulget (ielem, value)
c *** primary function:
                          get element nodal euler angles.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
C
С
        ielem
                (int,sc,in)
                                    - element number
     output arguments:
C
С
        eulget (int,func,out)
                                    - status of element.
                                      = 0 - element undefined
С
                                      > 0 - number of euler angle values
C
                                             returned
С
С
        value
                  (dp,ar(*),out)
                                    - element euler angles
С
                                  NOTE: For lower-ordered elements, rotations
C
                                          at centroid
                                        For higher-ordered elements, rotations
C
С
                                          at each corner node
                                        For layered shells, rotations at each
C
С
                                          corner node, plus layer rotation angle
С
                                          for each layer (real constant THETA)
                                        For layered solids, rotation angles at
C
                                          centroid, plus layer rotation angle
С
                                          for each layer (real constant THETA)
С
С
                                        For surface element, no euler angles
С
                                          are saved
```

3.7.47. Subroutine eulput (Storing an Element's Euler Angles)

```
*deck,eulput
     subroutine eulput (ielem,nval,value)
c *** primary function:
                          store nodal euler angles for an element.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
        ielem (int,sc,in)
                                     - element number
C
        nval
                                     - the total number of values
С
                  (int,sc,in)
                                       (3 * number of display nodes)
С
                                    - the euler angle values
        value
                 (dp,ar(nval),in)
C
С
     output arguments: none
C
                                  NOTE: For lower-ordered elements, rotations
С
                                          at centroid
                                        For higher-ordered elements, rotations
C
                                          at each corner node
С
С
                                        For layered shells, rotations at each
C
                                          corner node, plus layer rotation angle
                                          for each layer (real constant THETA)
С
С
                                        For layered solids, rotation angles at
                                          centroid, plus layer rotation angle
С
                                          for each layer (real constant THETA)
```

3.7.48. Subroutine euldel (Deleting an Element's Euler Angles)

3.7.49. Function efxiqr (Getting Information About Element Fluxes)

```
*deck,efxigr
     function efxiqr (ielem, key)
c *** primary function: get information about element fluxes
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
        ielem (int,sc,in)
С
                                  - element number (of 2011)
- key as to the information needed
                                    - element number (or zero, see below)
         key
C
                   (int,sc,in)
С
                  = 1 - return info about element fluxes
                              ielem > 0 - return number of fluxes on this
C
С
                                             element.
С
                                             (record length)
                                     = 0 - return maximum number of fluxes
С
С
                                             on any element
С
                                             (max record length)
                  = DB_NUMDEFINED - return the number of element fluxes defined
C
С
                                     in model
С
      output arguments:
         efxiqr (int,func,out)
С
                                    - the returned value of efxigr is based on
                                       setting of key
C
```

3.7.50. Function efxget (Getting an Element Flux)

```
*deck,efxget
      function efxget (ielem, value)
c *** primary function:
                        get element nodal fluxes.
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
        ielem (int,sc,in)
                                   - element number
С
С
     output arguments:
       efxget (int,func,out)
                                   - status of element.
C
С
                                     = 0 - element undefined
                                     > 0 - number of nodal fluxes returned
С
C
        value (dp,ar(*),out)
                                   - element nodal fluxes
                                   Note: If a coupled field, a set of fluxes is
C
                                          stored in the following order (as
C
С
                                          available): fluid, thermal,
                                          electric, magnetic
C
```

```
c *** mpg efxget<pagend<paberrwb,edgzzx,panavg,papres,paterr: get ele nd flx, B
```

3.7.51. Subroutine efxput (Storing an Element's Fluxes)

```
*deck,efxput
     subroutine efxput (ielem,nval,value)
c *** primary function:
                          store nodal fluxes at an element.
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
        ielem (int,sc,in)
                                    - element number
С
        nval
                  (int,sc,in)
                                    - the total number of values
С
                                      (ndir*nnod*nscalr)
С
                (dp,ar(nval),in) - the flux values
        value
C
С
     output arguments: none
С
                                   Note: If a coupled field, a set of fluxes is
                                           stored in the following order (as
С
                                          available): fluid, thermal,
C
                                          electric, magnetic
C
```

3.7.52. Subroutine efxdel (Deleting Element Fluxes)

3.7.53. Function elfiqr (Getting Information About Element Local Forces)

```
*deck,elfiqr
     function elfiqr (ielem, key)
c *** primary function: get information about elem local forces
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
C
        ielem (int,sc,in)
                                    - element number (or zero, see below)
С
                  (int,sc,in)
                                  - key as to the information needed
                  = 1 - return info about element local forces
C
                              ielem > 0 - return number of local forces on this
С
С
                                            element
                                            (record length)
C
                                    = 0 - return maximum number of local forces
С
С
                                            on any element
С
                                            (max record length)
```

```
c = DB_NUMDEFINED - return the number of element local forces
defined in model

c output arguments:
c elfiqr (int,func,out) - the returned value of elfiqr is based on
setting of key
```

3.7.54. Function elfget (Getting an Element Local Force)

```
*deck,elfget
     function elfget (ielem, value)
c *** primary function:
                        get element local nodal forces.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
       ielem (int,sc,in)
                                  - element number
С
    output arguments:
        elfget (int,func,out)
                                  - status of element.
                                    = 0 - element has no local nodal forces
С
                                    > 0 - number of nodal forces returned
        value
               (dp,ar(*),out) - element local nodal forces.
С
c *** mpg elfget<pagend<paberrwb,edgzzx,panavg,papres,paterr: get ele nd frc, F
```

3.7.55. Subroutine elfput (Storing an Element's Local Forces)

```
*deck,elfput
     subroutine elfput (ielem,nval,value)
c *** primary function: store element local nodal forces.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
     variable (typ,siz,intent) description
                 (int,sc,in)
                                 - element number
        ielem
C
С
        nval
                  (int,sc,in)
                                   - the total number of values
                                    NOTE: There may be a maximum of 3 sets of
С
C
                                    nodal forces in the record: static
С
                                    forces, inertia forces, and damping forces
        value
                 (dp,ar(nval),in) - element local nodal forces
C
     output arguments: none
```

3.7.56. Subroutine elfdel (Deleting Element Local Forces)

3.7.57. Function emniqr (Getting Information About Element Miscellaneous Non-summable Data)

```
*deck,emnigr
     function emniqr (ielem, key)
c *** primary function:
                          get information about element misc non-summable
                           data
С
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
                                    - element number (or zero, see below)
        ielem (int,sc,in)
С
         key
                  (int,sc,in)
                                    - key as to the information needed
С
                  = 1 - return info about element misc non-summed data
С
                              ielem > 0 - return number of data items on this
C
С
                                            element
С
                                            (record length)
                                    = 0 - return maximum number of data items
C
                                            on any element
С
                                            (max record length)
С
                  = DB_NUMDEFINED - return the number of element misc non-summed
С
                                     data items defined in model
С
С
      output arguments:
                                    - the returned value of emniqr is based on
С
         emniqr (int,func,out)
                                        setting of key
C
```

3.7.58. Function emnget (Getting an Element's Miscellaneous Non-summable Data)

```
*deck,emnget
     function emnget (ielem, value)
c *** primary function: get misc non-summable data.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
        ielem (int,sc,in)
                                   - element number
С
     output arguments:
                (int,func,out)
                                   - status of element.
С
         emnget
                                      = 0 - no non-summed misc data at this
C
                                            element
С
                                      > 0 - number of data items returned
С
        value
                 (dp,ar(*),out)
                                    - element misc non-summed data.
С
                                    NOTE: the contents of this record is element
C
                                       dependent. See NMISC on ETABLE command
C
```

3.7.59. Subroutine emnput (Storing an Element's Miscellaneous Non-summable Data)

```
*deck,emnput
     subroutine emnput (ielem,nval,value)
c *** primary function:
                         store misc. non-summable data for an element.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
С
       ielem (int,sc,in)
                                   - element number
                                   - the total number of values
С
        nval
                  (int,sc,in)
        value (dp,ar(nval),in) - the misc. non-summed data items
C
С
    output arguments: none
                                   NOTE: the contents of this record is element
С
                                      dependent. See NMISC on ETABLE command
С
```

3.7.60. Subroutine emndel (Deleting an Element's Miscellaneous Non-summable Data)

3.7.61. Function ecdiqr (Getting Information About Element Current Densities)

```
*deck,ecdiqr
     function ecdiqr (ielem, key)
c *** primary function: get information about element current densities
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
                                  element number (or zero, see below)key as to the information needed
        ielem (int,sc,in)
C
                   (int,sc,in)
C
         key
С
                  = 1 - return info about element current densities
                               ielem > 0 - return number of current densities on
С
                                             this element
С
С
                                             (record length)
                                     = 0 - return maximum number of current
C
С
                                             densities on any element
                                             (max record length)
С
С
                   = DB_NUMDEFINED - return the number of element current
С
                                      densities defined in model
c output arguments:
```

```
c ecdiqr (int,func,out) - the returned value of ecdiqr is based on setting of key
```

3.7.62. Function ecdget (Getting an Element Current Density)

```
*deck,ecdget
     function ecdget (ielem, value)
c *** primary function:
                        get calculated element current densities.
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
        ielem (int,sc,in)
                                  - element number
С
С
     output arguments:
       ecdget (int,func,out) - status of element.
C
С
                                    = 0 - element has no current densities
                                    > 0 - number of calculated element
С
С
                                           current densities
      value
               (dp,ar(*),out)
                                   - calculated element current densities.
С
                                   NOTE: current densities are in the order
С
С
                                         X, Y, Z
```

3.7.63. Subroutine ecdput (Storing an Element's Current Densities)

```
*deck,ecdput
     subroutine ecdput (ielem,nval,value)
c *** primary function: store calculated element current densities
c *** Notice - This file contains ANSYS Confidential information ***
С
     input arguments:
       ielem (int,sc,in)
                                   - element number
C
С
                                   - the total number of values
С
               (dp,ar(nval),in) - calculated element current densities.
С
     output arguments: none
                                   NOTE: current densities are in the order
C
C
                                         X, Y, Z
```

3.7.64. Subroutine ecddel (Deleting Element Current Densities)

3.7.65. Function enliqr (Getting Information About Element Nonlinear Tables)

```
*deck,enligr
     function enligr (ielem,key)
c *** primary function: get information about element nonlinear tables
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
        ielem
                (int,sc,in)
                                   - element number (or zero, see below)
C
С
                  (int,sc,in)
                                  - key as to the information needed
                  = 1 - return info about element nonlinear tables
C
                             ielem > 0 - return number of nonlinear tables for
С
С
                                            this element
                                            (record length)
C
                                    = 0 - return maximum number of nonlinear
                                           tables for any element
С
С
                                            (max record length)
                  = DB_NUMDEFINED - return the number of element nonlinear
С
                                    tables defined in model
C
С
     output arguments:
        enliqr (int,func,out)
                                   - the returned value of enliqr is based on
C
С
                                        setting of key
```

3.7.66. Function enlget (Getting Element Nonlinear Tables)

```
*deck,enlget
     function enlget (ielem, value)
c *** primary function: get element nonlinear tables.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
C
       ielem (int,sc,in)
                                   - element number
     output arguments:
C
        enlget (int,func,out)
                                   - status of element.
С
                                     = 0 - nonlinear tables undefined
С
                                     > 0 - number of nonlinear tables defined
C
        value (dp ,ar(n),out)
                                   - the element nonlinear tables.
С
                                   NOTE: Nonlinear data at each node are in the
С
                                          order SEPL, SRAT, HPRES, EPEQ, PSV,
С
                                          PLWK, and 4 spares
C
                                         For beam elements, the contents and
С
                                          number of information is element
С
С
                                          dependent. See NLIN on ETABLE
С
                                          command
```

3.7.67. Subroutine enlput (Storing an Element's Nonlinear Tables)

```
*deck,enlput
subroutine enlput (ielem,n,temp)
c *** primary function: store element nonlinear tables
```

```
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
С
        ielem
                (int,sc,in)
                                   - element number
                  (int,sc,in)
                                   - number of element nonlinear table values
C
        n
                 (dp ,ar(6),in)
                                   - element nonlinear table, etc.
С
C
     output arguments: none.
                                    NOTE: Nonlinear data at each node are in the
                                           order SEPL, SRAT, HPRES, EPEQ, PSV,
С
С
                                           PLWK, and 4 spares
С
                                          For beam elements, the contents and
                                           number of information is element
C
С
                                          dependent. See NLIN on ETABLE
С
```

3.7.68. Subroutine enIdel (Deleting Element Nonlinear Tables)

3.7.69. Function ehciqr (Getting Information About Calculated Element Heat Generations)

```
*deck,ehciqr
     function ehciqr (ielem, key)
c *** primary function: get information about calculated elem heat generations
c *** Notice - This file contains ANSYS Confidential information ***
      input arguments:
С
С
        ielem
                (int,sc,in)
                                    - element number (or zero, see below)
                                    - key as to the information needed
                  (int,sc,in)
C
        key
С
                  = 1 - return info about calculated element heat gens
                          for ielem > 0 - return number of heat gens for
С
С
                                            this element
С
                                            (record length)
                                    = 0 - return maximum number of heat gens
C
С
                                            for any element
                                            (max record length)
С
                  = DB_NUMDEFINED - return the number of calculated element heat
С
                                     generations defined in model
С
С
      output arguments:
        ehciqr (int,func,out)
                                    - the returned value of ehciqr is based on
С
                                        setting of key
C
```

3.7.70. Function ehcget (Getting a Calculated Element Heat Generation)

```
*deck,ehcget
     function ehcget (ielem, value)
c *** primary function: get calculated element heat generations.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
       ielem (int,sc,in)

    element number

С
    output arguments:
C
С
       ehcget (int,func,out) - status of element.
                                   = 0 - element undefined
С
                                   > 0 - number of calculated element
С
                                         heat generations
С
    value (dp,ar(*),out) - calculated element heat generations.
```

3.7.71. Subroutine ehcput (Storing an Element's Calculated Heat Generations)

3.7.72. Subroutine ehcdel (Deleting Element Calculated Heat Generations)

Chapter 4: Subroutines for Your Convenience

This chapter describes routines available to you for use in programming. Using these routines isn't required, but may make your life easier. These routines include a set of general routines that perform utility-type functions, a set of routines supporting vector functions, a set of routines supporting matrix functions, and routines supporting message processing options.

The following topics are discussed in this chapter:

- 4.1. Input and Output Abbreviations
- 4.2. General Subroutines
- 4.3. Vector Functions
- 4.4. Matrix Subroutines

4.1. Input and Output Abbreviations

The descriptions of inputs and outputs for the routines discussed in this chapter use the following abbreviations:

• Argument *type* is one of the following:

int - integer

dp - double precision

log - logical

chr - character

dcp - double precision complex

• Argument size is one of the following:

sc - scalar variable

ar(n) - array variable of length n

func - functional return value

• Argument intent is one of the following:

in - input argument

out - output argument

inout - both an input and an output argument

4.2. General Subroutines

The following general subroutines are available for your convenience:

- 4.2.1. Subroutine dptoch (Retrieve Eight Characters From a Double Precision Variable)
- 4.2.2. Function ppingr (Obtain Information About Threads)
- 4.2.3. Function pplock (Locking a Thread in Shared Memory)
- 4.2.4. Function ppunlock (Unlocking a Thread in Shared Memory)
- 4.2.5. Function ppproc (Get the Active Thread Index)
- 4.2.6. Function wringr (Obtain Information About Output)
- 4.2.7. Subroutine eringr (Obtaining Information from the Errors Common)
- 4.2.8. Subroutine erhandler (Displaying Program Errors)
- 4.2.9. Subroutine intrp (Doing Single Interpolation)
- 4.2.10. Subroutine tranx3 (Processing Geometry for 3-D Line Elements)
- 4.2.11. Subroutine systop (Stopping a Program Run)

4.2.1. Subroutine dptoch (Retrieve Eight Characters From a Double Precision Variable)

```
*deck,dptoch
subroutine dptoch (dp8,ch8)

c *** primary function: retreive 8 characters from a dp variable

c *** Notice - This file contains ANSYS Confidential information ***

c !!! NOTICE to progammers: this routine does not convert from a !!!

c !!! machine-independent format! Use dpexttoch if this dp word !!!

c !!! came from a common or non-char database record !!!

c input arguments:

c dp8 (dp,sc,in) - dp variable containing characters

c output arguments:

c ch8 (ch*8,sc,out) - characters retreived from the dp word
```

4.2.2. Function ppingr (Obtain Information About Threads)

```
*deck,ppinqr
function ppinqr (key)

c primary function: Get information from pplib (parallel library)

c keywords: subroutine to inquire from parallel library

c object/library:
c current - pplib

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
c key (int,sc,in) - 0, Get ppNprocs
c (if >1, parallel allowed)
c 1, Get ppOff
```

```
C
                                         (0,pp active 1, inactive)
С
                                     2, Get ppMaxProc
                                         (max processors to be used)
C
                                     3, Get ppNumThreads
C
С
                                         (number of threads in set)
                                     4, Get ppThdRunning
С
                                         (either 0 or the number of threads running)
С
С
                                     5, Get ppDebug
                                         (debug level value)
C
                                     7, -1, parallel not started
                                         0, parallel started, threads not
C
С
                                        n, number of threads running
С
                                    10, Get ppNprocOrig
                                    12, Get ppInitialize
C
С
                                    13, Get ppRunLevel
С
                                    22, Get ppLockCheck
                                    23, Get ppHybrid
C
С
   output arguments:
      ppingr (int,sc,out)
                                 - The value of the referenced variable
```

For information about using this function, see Sharing Data Between User Routines (p. 137).

4.2.3. Function pplock (Locking a Thread in Shared Memory)

```
*deck,pplock
subroutine pplock (ilock)
c ******** set the lock ilock ********
```

For information about using this function, see Sharing Data Between User Routines (p. 137).

4.2.4. Function ppunlock (Unlocking a Thread in Shared Memory)

```
*deck,ppunlock
subroutine ppunlock(ilock)
c ********* clear the lock ilock *********
```

For information about using this function, see Sharing Data Between User Routines (p. 137).

4.2.5. Function ppproc (Get the Active Thread Index)

```
*deck,ppproc
    function ppproc ()
c *** primary function: return unique thread number (0:ppNprocs-1)
c *** Notice - This file contains ANSYS Confidential information ***
c output arguments:
c ppproc (int,sc,out) - the thread number for this process
```

For information about using this function, see Sharing Data Between User Routines (p. 137).

4.2.6. Function wringr (Obtain Information About Output)

```
*deck,wringr
     function wringr (key)
c *** primary function:
                         obtain information about output
c *** Notice - This file contains ANSYS Confidential information ***
c --- caution: the following variables are "saved/resumed".
              key=WR_COLINTER thru WR_SUPCOLMAX in "wringr/wrinfo"
C ---
               (data for "/fmt,/page,/header" commands).
               note that the whole common cannot be "saved/resumed". cwa
C ---
     typ=int,dp,log,chr,dcp siz=sc,ar(n),func intent=in,out,inout
С
  input arguments:
     variable (typ,siz,intent) description
С
                                                                      wrcom name
C
     key
          (int,sc,in)
                   = WR_PRINT - print flag (kprint)
                                                                          prtkey
С
                       wringr = 0 - no output
С
                            = 1 - print
C
                  = WR_OUTPUT - current output unit number(iott) outfil
С
                  = WR_MASTEROUT - master output file
C
С
                  = WR COLINTER
                                   - interactive columns per page
                  = WR_COLBATCH - batch columns per page
                                                                     batcol
С
                  = WR_LINEINTER - interactive lines per page
                                                                      intlin
C
                  = WR_LINEBATCH - batch lines per page
С
                                                                     batlin
                  = WR_COMMASEP - 1 for comma separated output
= WR_CHARITEM - characters per output item
                                                                     CommaSep
С
С
                                                                      chrper
                  = WR_CHARDECIMAL - characters past decimal
С
                                                                       chrdec
                  = WR_CHARINTEGER - characters in leading integer
                                                                      chrint
С
                  = WR_CHARTYPE
                                                                       chrtyp
С
                        wringr = 1 - using E format in output
C
                              = 2 - using F format in output
C
                               = 3 - using G format in output
С
С
                  = WR_SUPTITLE - tlabel supress key
                                                                       keyhed
                  = WR SUPSUBTITLE - subtitle supress key
                                                                       keytit
C
С
                  = WR_SUPLSITER - ls,iter id supress key
                                                                       keyid
                  = WR_NOTELINE - note line supress key
С
                                                                      keynot
С
                  = WR_SUPCOLHEADER - column header supress key
                                                                       keylab
                  = WR_SUPCOLMAX - column maximum supress key
= WR_LISTOPT - ListOpt from /output command
                                                                       keysum
С
                                                                      ListOpt
C
c output arguments:
     wringr (int, func, out) - the value corresponding to key
```

4.2.7. Subroutine eringr (Obtaining Information from the Errors Common)

```
*deck,eringr
     function eringr (key)
c *** primary function:
                        obtain information from errors common
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
     key (int,sc,in)
                                - item to be returned
C
                                  1=keyerr, 2=errfil, 3=numnot, 4=numwrn,
С
                                   5=numerr, 6=numfat,
                                                         7=maxmsg, 8=lvlerr,
C
                                   9=mxpcmd, 10=nercmd, 11=nertim, 12=nomore,
С
                                   13=eropen,14=ikserr, 15=kystat,16=mxr4r5,
С
C
                                                        19=opterr, 20=flowrn,
С
                                            22=noreport, 23=pdserr, 24=mxpcmdw
C
                                   25=kystop,26=icloads, 27=ifkey,
C
                                   28=intrupt
```

```
С
c ---- below definitions copied from errcom 7/92 for user information
C
                             *** key number= .....
C
С
                     (see ansysdef for parameter definitions)
C
С
                                                                  (ER_ERRORFLAG)
co keyerr - master error flag
co errfil - errors file unit number
                                                                  (ER ERRORFILE)
co numnot - total number of notes displayed
                                                                  (ER_NUMNOTE)
co numwrn - total number of warnings displayed
                                                                  (ER_NUMWARNING)
co numerr - total number of errors displayed co numfat - total number of fatals displayed
                                                                  (ER_NUMERROR)
                                                                  (ER_NUMFATAL)
co maxmsg - max allowed number of displayed messages before abort(ER_MAXMESSAGE)
co lvlerr - used basicly in solution (from cnvr command.)
                                                                  (ER_ERRORLEVEL)
             -1=do not set keyerr for notes/errors/warnings.
CO
             -2=same as -1 but do not display message either.
CO
co mxpcmd - maximum number of messages allowed per command
                                                                  (ER_MAXCOMMAND)
co nercmd - number of messages displayed for any one command
                                                                  (ER_NUMCOMMAND)
co nertim - key as to how message cleared from u/i pop-up
                                                                  (ER_UICLEAR)
            (as per rsg/pft 5/1/92 - only for "info" calls
             -1=message is timed before removal
CO
              0=message needs pick or keyboard before removal
CO
              1=message stays up untill replaced by another message
CO
           display any more messages
                                                                  (ER NOMOREMSG)
co nomore
CO
             0=display messages
             1=display discontinue message and stop displaying
CO
co eropen - 0=errors file is closed
                                                                  (ER FILEOPEN)
            1=errors file is opened
CO
co ikserr - 0=if interactive do not set keyerr
                                                                  (ER_INTERERROR)
          - 1=if interactive set keyerr (used by mesher and tessalation)
C
co kystat - flag to bypass keyopt tests in the elcxx routines
                                                                  (ER_KEYOPTTEST)
C
            associated with status/panel info inquiries.
             0=do not bypass keyopt tests
С
С
             1=perform all keyopt tests
            also flag to bypass setting of _STATUS upon resume
C
co mxr4r5 - mixed rev4-rev5 input logic (*do,*if,*go,*if-go)
                                                                  (ER MIXEDREV)
             (used in chkmix called from rdmac)
С
             1=rev5 found (*do,*fi-then-*endif)
С
C
             2=rev4 found (*go,:xxx,*if,...,:xxx)
             3=warning printed. do not issue any more.
С
co mshkey - cpu intensive meshing etc. this will cause
                                                                  (ER_MESHING)
            "nertim (11)" to be set to -1 for "notes", 1 for "warnings",
С
С
            and 0 for "errors". checking of this key is done in "anserr".
             0=not meshing or cpu intensive
С
             1=yes, meshing or cpu intensive
co syerro - systop error code. read by anserr if set.
co opterr - 0=no error in main ansys during opt looping
                                                                  (ER_OPTLOOPING)
           1=an error has happened in main ansys during opt looping
co flowrn - flag used by "floqa" as to list floqa.ans
            0=list "floqa.ans"
C
            1="floqa.ans" has been listed. do not list again.
co noreport- used in GUI for turning off errors due to strsub calls (22)
            0=process errors as usual
C
            1=do NOT report errors
C
co pdserr - 0=no error in main ansys during pds looping
                                                                  (ER_PDSLOOPING)
            1=an error has happened in main ansys during pds looping
co mxpcmdw- number of messages written to file.err for any one
            command
CO
С
            0=write all errors to file.err
            1=only write displayed errors to file.err
co icloads - key to forbid the iclist command from listing solution
             data instead of the input data.
С
            0=iclist is OK
C
           1=do not permit iclist
           - key on whether or not to abort during /input on error
                                                                        (27)
co ifkey
            0=do not abort
C
            1=abort
co intrupt - interrupt button, so executable returns no error
                                                                  (ER INTERRUPT)
co espare - spare integer variables
```

```
c
c --- end of information from errcom
c
c output arguments:
c erinqr (int,sc,out) - value corresponding to key
c
c *** mpg erinqr < el117,el115,el126,el109,el53,el96,el97,edg?: get error stat
c</pre>
```

4.2.8. Subroutine erhandler (Displaying Program Errors)

```
*deck,erhandler
      subroutine erhandler (filein, msgid, msglvl, lngstrng, dperr, cherr)
c *** primary function:
                           Display ANSYS error messages
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
C
                In order to make life for vismg easier,
C
                do NOT use variables for the first four arguments
C
С
С
      filein (ch*40,sc,in)
                                  - Filename used for character portion of
                                    message ID (this is the file name of the
C
                                    file which contains the source for this
С
С
                                    routine)
C
C
                                    if 'ErrorMessageProbe', then error was
                                      generated on another processor (distributed
C
                                      ANSYS). In that case, dperr contains the
С
                                      message already made ASCII and expanded
С
С
                                  - Numeric portion of the message ID
С
      msgid
               (int,sc,in)
С
                                    1 - 9999, unique for each erhandler
                                    call in the FILE. Recommend using
С
С
                                    a sequence, similar to format conventions,
                                    i.e., 5000, 5010, 5020
С
                                    if filein='ErrorMessageProbe', this is the
C
                                      CPU # that originally generated the error
С
С
      msglvl
               (int,sc,in)
                                  - level of error (same as lngerr)
                                    0=no label (used for u/i pop-ups)
C
С
                                   -1=no label (used for u/i pop-ups) timed
                                      as a note message
С
С
                                    1=note, 2=warning, 3=error, 4=fatal
С
                                                      -3=error w/tech supp note
                                                      -4=fatal w/tech supp note
C
                                       (see lngerr.F for text of tech supp note)
С
С
      lngstrng (ch*(*),sc,in)
                                  - error message to display. use keywords
                                    of %i %g %c %/ for formating (same as
C
С
                                    lngerr)
С
      dperr
               (dp,ar(*),in)
                                  - vector of data to display. contains both
                                    integer and double precision data.
C
С
                                    (same as lngerr)
                                      if filein='ErrorMessageProbe', dperr
C
С
                                     contains the unpacked message and lngstrng
С
                                     and cherr are ignored
               (ch*(*),ar(*),in) - vector of character data to display
C
      cherr
                                    max length of character data is 32
С
С
                                    characters
```

4.2.9. Subroutine intrp (Doing Single Interpolation)

```
*deck,intrp
     subroutine intrp (klog,kppx,kstpz,xval,ax,ay,yval,nmax,kyoff)
c *** primary function: **** subroutine for single interpolation ****
                        (if double interpolation is needed, see intrpt)
С
С
c *** Notice - This file contains ANSYS Confidential information ***
C
С
      typ=int,dp,log,chr,dcp siz=sc,ar(n),func
С
  input arguments:
C
С
   variable (typ,siz,intent)
                                  description
                                  - interpolation type
С
     klog
               (int,sc,in)
                                    = 0 - use linear interpolation
C
                                    = 1 - use log-log interpolation
С
                                      -- note: there is no option yet for
С
                                                 lin-log or log-lin
C
С
      kppx
                (int,sc,in)
                                  - X value end of table signal
                                    = 0 - a repeated x-value will signal the end
C
                                          of the table
C
С
                                    = 1 - a repeated x-value will not signal the
                                          end of the table
С
С
                                           (only known use = c evaluation)
                                  - Y value end of table signal
С
      kstpz
                (int,sc,in)
                                    = 0 - a yval of zero will not signal the end
C
                                          of the table (e.g. stress fitting)
С
                                    = 1 - a yval of zero will signal the end of
С
С
                                           the table (in general, material
С
                                          properties (exception: alpx))
С
С
                          NOTE: the end of the table will be signaled thru
                                either of the above conditions, or more
C
                                commonly, that nmax values have been processed,
C
                                or that the present x table entry is less than
С
                                the previous one (ax(i) .lt. ax(i-1)).
С
                                evaluations done after the end of the table are
C
С
                                evaluated as if they were at the end of the
                                table. similarly, evaluations done before the
C
С
                                beginning of the table are done as if they were
                                done at the beginning of the table.
С
C
      xval
               (dp,sc,in)
                                  - value of x with which to go into the table
С
С
      ax
               (dp,ar(*),in)
                                  - table of x values, in ascending order
               (dp,ar(*),in)
                                  - table of y values
С
      ay
               (int,sc,in)
                                  - maximum table size allowed
С
      nmax
С
  output arguments:
C
     yval
               (dp,sc,out)
                                  - value of y which comes back from the table
С
     kyoff
                                  - xval status flag
               (int,sc,out)
С
                                    = 0 - xval in x range
C
                                    = 1 - xval less than minimum x
С
                                    = 2 - xval greater than maximum x
C
C
```

4.2.10. Subroutine tranx3 (Processing Geometry for 3-D Line Elements)

```
c nnod (int,sc,in) - number of nodes (2 or 3)
c xyz (dp,ar(nx,*),in) - coordinates (x,y,z down)
c nx (int,sc,in) - row dimension of xyz array
c
c output arguments:
c tr (dp,ar(3,3),in) - transformation matrix
c
```

4.2.11. Subroutine systop (Stopping a Program Run)

```
*deck,systop
     subroutine systop (icode)
c *** primary function: stop an ansys run
c *** secondary functions: pass an error code to the system
c *** Notice - This file contains ANSYS Confidential information ***
C
c input arguments:
С
     icode (int,sc,in)
                                 - stop error code (0<icode<127)
С
                                    0 - normal exit
                                    1 - stack overflow error
С
                                    2 - stack level overflow
С
                                    3 - stack pop below zero
С
С
                                    4 - names do not match in stkpxp
                                    5 - command line argument error
C
С
                                    6 - unused (was: accounting file error)
С
                                    7 - licensing failure
                                    8 - indicated error or end-of-run
C
С
                                   11 - error in user routine
                                   12 - macro stop command
C
                                   13 - job already running
С
                                   14 - untrapped xox error
С
                                   15 - anserr fatal error
C
                                   16 - possible full disk
С
С
                                   17 - possible corrupted or missing file
С
                                   18 - Error in VM routines (corrupt db?)
С
                                   21 - unauthorized code section entered
                                   25 - unable to open x11 server
С
                                   30 - quit signal
C
                                   31 - failure to get signal in max time
С
С
                                           (syhold)
                                  >32 - system dependent error
C
                                   35 - fatal error on another process
С
                                         (distributed ANSYS)
C
  output arguments: none
```

4.3. Vector Functions

4.3.1. Function vdot (Computing the Dot Product of Two Vectors)

```
*deck,vdot
     function vdot (v1,v2,n)
c *** primary function: compute dot product of vectors v1 and v2
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
c v1 (dp,ar(n),in) - vector v1
```

4.3.2. Function vidot (Computing the Dot Product of Two Vectors with Increments)

```
*deck,vidot
function vidot (v1,inc1,v2,inc2,n)
c *** primary function: compute the dot product of vectors v1 and v2
c *** Notice - This file contains ANSYS Confidential information ***
c
---- incl and inc2 must be positive!
c
```

4.3.3. Function vsum (Summing Vector Components)

4.3.4. Function vmax (Retrieving the Maximum Vector Value at a Given Location)

```
*deck,vmax
function vmax (v,n,locmax)
c *** primary function: get the biggest value and location in v
c *** Notice - This file contains ANSYS Confidential information ***
```

4.3.5. Function lastv (Retrieving the Position of the Last Nonzero Term in a Double Precision Vector)

```
*deck,lastv
function lastv (v,n)
c ******* find position of last non-zero term in a d.p. vector *******
```

4.3.6. Function izero (Setting an Integer Vector to Zero)

```
*deck,IZERO
subroutine IZERO (ivect,n)
c ******** set an integer vector to zero *********
```

4.3.7. Function imove (Assigning Equal Values to Two Integer Vectors)

```
*deck,imove subroutine imove (i1,i2,n) c ******* move a vector from one to another ********
```

4.3.8. Subroutine vzero (Initializing a Vector to Zero)

4.3.9. Subroutine vmove (Moving One Vector into Another)

```
*deck,vmove
subroutine vmove (v1,v2,n)

c *** primary function: copy v1 vector into another vector

c *** Notice - This file contains ANSYS Confidential information ***
```

4.3.10. Subroutine vimove (Moving One Vector into Another Incrementally)

4.3.11. Subroutine vinit (Assigning a Scalar Constant to a Vector)

```
*deck, vinit
    subroutine vinit (v,n,const)

c *** primary function: initialize a vector to a constant

c *** Notice - This file contains ANSYS Confidential information ***
```

4.3.12. Subroutine viinit (Assigning a Scalar Constant to a Vector Incrementally)

```
*deck,viinit
```

```
subroutine viinit (v,inc,n,const)
c *** primary function: set the components of vector v to const by increments
c *** Notice - This file contains ANSYS Confidential information ***
```

4.3.13. Subroutine vapb (Setting a Vector to Sum of Two vectors)

4.3.14. Subroutine vapb1 (Combining Two Vectors in One)

```
*deck,vapb1
    subroutine vapb1 (a,b,n)
c *** primary function: add vector b to vector a and store in vector a
```

4.3.15. Subroutine vapcb1 (Multiplying a Vector to a Constant)

4.3.16. Subroutine vamb (Gets a Third Vector by Subtracting One Vector from Another)

4.3.17. Subroutine vamb1 (Subtracting One Vector from Another)

4.3.18. Subroutine vmult (Multiplying a Vector by a Constant)

```
*deck,vmult
subroutine vmult (v1,v2,n,const)
c *** primary function: multiply a vector by a constant
```

```
c *** Notice - This file contains ANSYS Confidential information ***
```

4.3.19. Subroutine vmult1 (Multiplying a Vector by a Constant)

4.3.20. Subroutine vcross (Defining a Vector via a Cross Product)

4.3.21. Subroutine vnorme (Normalizing a Three-Component Vector)

```
*deck, vnorme
     subroutine vnorme (iel,v)
c primary function: normalize a vector to unit length
c this routine is to be called only from the elements.
                                                       it is only
c for a three component vector(i.e. processing geometry).
  this routine also differs from vnorm in that an error message is called
c if the vector length is zero.
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
    iel (int,sc,inout) - element number
C
             (dp,ar(3),inout) - vector to be normalized
c output arguments:
    iel (int,sc,inout) - if 0, vector has zero length
              (dp,ar(3),inout) - normalized vector
C
```

4.3.22. Subroutine vnorm (Normalizing a Vector to Unit Length)

```
*deck,vnorm
```

```
subroutine vnorm (v,n)
c *** primary function: normalize a vector to unit length

c *** Notice - This file contains ANSYS Confidential information ***

c input arguments:
    v     (dp,ar(n),inout) - vector v
    n     (int,sc,inout) - dimension length of vector v

c output arguments:
    v     (dp,ar(n),inout) - normalized vector v

c n     (int,sc,inout) - n = 0 if error in operation
```

4.3.23. Function ndgxyz (Getting the X,Y,Z Vector for a Node)

```
*deck,ndgxyz
     function ndgxyz (node,xyz)
c *** primary function: get x,y,z vector for a node.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
С
       node (int,sc,in)
                                - node number for operation.
С
С
     output arguments:
       ndgxyz (int,sc,out)
                                 - status of node.
                                     0=node is undefined.
C
                                     -1=node is unselected.
                                     1=node is selected.
С
    xyz (dp,ar(3),out)
                                 - vector containing x,y,z
С
```

4.3.24. Subroutine ndpxyz (Storing X,Y,Z for a Node)

```
*deck,ndpxyz
     subroutine ndpxyz (node,xyz)
c *** primary function: store x,y,z vector for a node.
c *** Notice - This file contains ANSYS Confidential information ***
     input arguments:
C
С
       node (int,sc,in)
                                  - node number for operation.
                               vector containing x,y,z
               (dp,ar(3),in)
C
                                    (vector should be in global system)
С
    output arguments: none
C
```

4.4. Matrix Subroutines

4.4.1. Subroutine maxv (Multiplying a Vector by a Matrix)

```
*deck,maxv
subroutine maxv (a,v,w, nr,nc)
c *** primary function: multiply a matrix by a vector
```

```
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
С
             (dp,ar(nr,*),in) - matrix a
              (dp,ar(*),in) - vector v
(int,sc,in) - number of rows in matrix a
C
     V
              (int,sc,in)
С
     nr
                               - number of columns to multiply in matrix a
С
     nc
              (int,sc,in)
c output arguments:
  w (dp,ar(*),out) - product vector w
C
С
c *** mpg w = A v : A(nr,nc) : matrix vector product
```

4.4.2. Subroutine maxv1 (Multiplying a Vector by a Matrix)

```
*deck,maxv1
     subroutine maxv1 (a,v, nr,nc)
c *** primary function: multiply a vector by a matrix
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
     a (dp,ar(nr,nc),in) - matrix a
C
С
              (dp,ar(nc),inout) - vector v
С
     nr
             (int,sc,in) - number of rows in matrix a
                                 *** nr limited to 60 ***
C
С
             (int,sc,in)
                               - number of columns to multiply in matrix a
 output arguments:
C
             (dp,ar(nr),inout) - product, stored in vector v
C
c *** mpg v = A v : A(nr,nc) : matrix vector product, max 60 rows
```

4.4.3. Subroutine matxv (Multiplying a Vector by a Full Transposed Matrix)

```
*deck,matxv
     subroutine matxv (a,v,w, nr,nc)
c *** primary function: multiply vector by full transposed matrix
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
          (dp,ar(nr,*),in) - matrix a (first dimension must = nr)
С
                                 - vector v (nv must be greater or equal
              (dp,ar(nv),in)
C
С
                                 - first dimension and number of active
С
    nr
              (int,sc,in)
                                      rows of the untransposed matrix a
С
С
                                       (also the number of active rows
                                       of vector v)
С
С
             (int,sc,in)
                                 - number of columns of the untransposed
С
                                      matrix a
                                      (also the number of computed items
C
С
                                      in the product vector w)
С
                                      if negative, accumulate
C
              (dp,ar(na,*),out) - product vector w
C
c *** mpg A(nr,nc) : matrix transpose vector product
```

```
c 	 w = A + v : if nr > 0
c 	 w = w + A + v : if nr < 0
c
```

4.4.4. Subroutine matxv1 (Multiplying a Vector by a Full Transposed Matrix)

```
*deck,matxv1
     subroutine matxv1 (a,v, nr,nc)
c *** primary function: multiply vector by full transposed matrix
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
     a (dp,ar(nr,*),in)
                                 - matrix a
C
              (dp,ar(nr),inout) - vector v
С
              (int,sc,in)
                                - number of rows in matrix (un-transposed)
С
     nr
                                - number of columns in matrix (un-transposed)
    nc
             (int,sc,in)
С
                                  *** nc limited to 60 ***
c output arguments:
              (dp,ar(nc),inout) - product, stored in vector v
С
c *** mpg A(nr,nc) : matrix transpose vector product
         v = A + v : max 60 nc
```

4.4.5. Subroutine matxb (Transposing a matrix)

```
*deck,matxb
     subroutine matxb (a,b,c, na,nb,nc, n1,n2,n3)
c *** primary function: (a)t * (b) = (c) t means transpose
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
С
          (dp,ar(na,*),in) - matrix a
             (dp,ar(nb,*),in) - matrix b
С
     b
             (int,sc,in) - number of rows in matrix b
С
     na
С
     nb
    nc
             (int,sc,in)
                               - number of rows in matrix c
С
С
    n1
             (int,sc,in)
                               - number of rows in matrix c to fill
C
    n2
            (int,sc,in)
                               - number of columns in matrix c to fill
    n3
C
             (int,sc,in)
                               - number of rows in matrix a and
                                 number of rows of matrix b
С
                                 to work with (the two need
С
                                 to be the same for the inner product)
C
                                 if n3 is negative, accumulate results in c
С
c output arguments:
             (dp,ar(nc,*),out) - product matrix c
c *** mpg C = A+ B if n3 > 0
        C = C + A + B
                        if n3 < 0
         A(na,*) B(nb,*) C(nc,*) C:minor n1 * n2 n3: dot length
C
С
```

4.4.6. Subroutine maat (Changing a Matrix Value via Addition, Multiplication, and Transposition)

```
*deck, maat
      subroutine maat(a,c, nc,n, con)
c primary function: does con*a*at and sums the result onto c (a is a vector)
c *** Notice - This file contains ANSYS Confidential information ***
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
С
  input arguments:
             (dp,ar(*),in)
                                  - vector to be multiplied by itself to
C
     a
                                   generate an nxn square matrix
                                     (a by a-transposed)
C
С
             (dp,ar(nc,*),inout) - matrix to be accumulated onto
              (int,sc,in)
                                 - number of rows in the c matrix
С
     nc
                                - size of square matrix
    n
              (int,sc,in)
C
                                - multiplier on above square matrix
С
    con
             (dp,sc,in)
c output arguments:
     c (dp,ar(nc,*),inout) - matrix to be accumulated onto
С
С
                                    only the lower triangular matrix is done
С
         Note: this routine is usually followed by matsym,
                                             to do the complete matrix
С
С
```

4.4.7. Subroutine matba (Updating Matrix Value via Transposition, Multiplications, and Addition)

```
*deck.matba
      subroutine matba (a,b,c,na,nb,nc,n1,n2,work,con)
c primary function: does con(at*b*a) and sums the result
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
         (dp,ar(na,*),in)
                                - matrix a
C
     а
     b
              (dp,ar(nb,*),in)
                               - matrix b (must be square,
С
                                        and maximum dimension is (15,15)
С
             (dp,ar(nc,*),inout)- matrix c (see output)
С
             (int,sc,in) - number of rows in matrix a
С
    na
    nb
             (int,sc,in)
                                - number of rows in matrix b
C
                                - number of rows in matrix c
С
     nc
              (int,sc,in)
С
     n1
             (int,sc,in)
                                - number of rows in matrix a and
                                  number of rows of matrix b
С
                                  to work with (the two need
С
С
                                  to be the same for the inner product)
С
     n2
              (int,sc,in)
                                 - number of columns in matrix c to fill
С
              (dp,sc,in)
                                 - multiplier on product added to sum
     con
 output arguments:
С
С
              (dp,ar(nc,*),inout)-c=c+con*at*b*a
              (dp,ar(n2,*),out) - at*b (this byproduct is occasionally useful)
С
c *** mpg C = C + con A+ B A A(na,*) B(nb,*) C(nc,*) C:minor n1 * n2
     see matbabd for block diagonal
C
С
```

4.4.8. Subroutine matsym (Filling the Upper Triangle from the Lower Triangle)

```
*deck,matsym
     subroutine matsym (a,nd,n)
c primary function: fill upper triangle from lower triangle
c *** Notice - This file contains ANSYS Confidential information ***
     typ=int,dp,log,chr,dcp siz=sc,ar(n) intent=in,out,inout
c input arguments:
             (dp,ar(nd,*),inout) - matrix to have its lower triangular part
C
С
                                    copied to its upper triangular part
                                 - number of rows of the a matrix
              (int.sc.in)
С
     nd
              (int,sc,in)
                                 - size of matrix to be processed
C
c output arguments:
             (dp,ar(nd,*),inout) - matrix that has its lower triangular part
C
С
                                     copied to its upper triangular part
C
```

4.4.9. Subroutine mctac (Transposing a symmetric matrix)

```
*deck, mctac
     subroutine mctac (a,na,c,nc,nold,nnew)
   **** function: do a = c(transpose) * a * c , where a is symmetric **
c *** Notice - This file contains ANSYS Confidential information ***
c input arguments:
С
        (dp,ar(na,na),inout) matrix to be pre and post multiplied
С
                                     (part operated on must be
                                    square(nold x nold) and symmetric)
C
                                   first dimension of the a matrix
С
              (dp,ar(nc,nnew),in) matrix to pre and post multiply a by
С
С
                                   (part used may be rectangular(nold x nnew))
С
     nc
              (int,sc,in)
                                   first dimension of the c matrix
                                   size of part of 'A' matrix that is
     nold
             (int,sc,in)
C
                                   to be processed(input size). maximum = 64
С
             (int,sc,in)
С
     nnew
                                   size of part of 'A' matrix that
                                    results from this operation(output size).
C
С
C
 output arguments:
С
           (dp,ar(na,na),inout) resulting matrix
                                     (still square(nnew x nnew) and symmetric).
C
```

4.4.10. Subroutine tran (Transposing a matrix)

```
C
     nz
              (int,sc,in)
                                  - dimensioned size of zs matrix
                                  - dimensioned size of tr matrix
С
     ntr
              (int,sc,in)
                                  - number of rows of zs matrix to transform
              (int,sc,in)
C
     nrow
     irot
              (int,sc,in)
                                  - block size to transform(size of tr matrix)
C
c output arguments:
     variable (typ,siz,intent)
С
                                 description
              (dp,ar(nz,nz),inout) - transformed matrix
```

4.4.11. Subroutine symeqn (Solving Simultaneous Linear Equations)

```
*deck,symeqn
     function symeqn (a,nd,n,nc,defFlag)
c primary function: solve a set of simultaneous linear equations
c secondary functions: invert a matrix
С
           NOTE: this routine assumes that the matrix to be solved or
                  inverted is positive or negative definite. This routine
                  also assumes that the diagonals are all non-zero. If
C
                  this assumption is not true, use isimeq.F.
c *** Notice - This file contains ANSYS Confidential information ***
С
  input arguments:
C
     variable (typ,siz,intent)
                                   description
C
C
              (dp,ar(nd,*),inout) - matrix to be solved or inverted
                                       second dimension must be at least:
С
С
                                                                 n + abs(nc)
                                   - first dimension of the a matrix
               (int,sc,in)
С
               (int.sc.in)
                                   - number of equations
С
     n
                                    - number of additional columns.
С
               (int,sc,in)
С
                                        if nc = +n or -n, invert n \times n matrix and
                                        put result in the n+1 to 2xn columns.
С
С
                                        if nc is 0 or negative, nc will be reset to
С
                                        n and then symeqn will set up identity
                                        matrix after the input matrix, where the
C
                                        result of the inversion will be put.
C
С
                                        if nc is positive and less than n, do a
                                        partial inversion. see example 1 below.
C
С
     defFlag (int,sc,in)
                                    - flag indicating that incoming matrix MUST be:
                                        -1 - negative definite
C
С
                                         0 - positive or negative definite
С
                                         1 - positive definite
C
С
  output arguments:
С
     variable (typ,siz,intent)
                                   description
      symeqn (in,sc,out)
                                    - 0 - non-singular matrix
C
                                     1 - singular matrix
С
С
                                     2 - near-singular matrix
               (dp,ar(nd,*),inout) - results or inverted matrix.
C
                                        starts in column n+1.
С
                                       note: original information is destroyed.
C
C
    example 1:
                 Solve three simultaneous linear equations:
                       i = symeqn (a(1,1),3,3,1)
C
С
                    calling routine has a dimensioned as a(3,4)
                    each equation has its {\tt 3} coefficents in the first {\tt 3} columns,
С
C
                     and the constant term is in the fourth column.
                    solution is in fourth column.
С
C
    example 2:
                 Invert a 3x3 matrix:
C
                       i = symeqn (a(1,1),3,3,-3)
C
C
                    calling routine has a dimensioned as a(3,6)
                    input matrix was input in first 3 columns
С
```

c output matrix in ouput in last 3 columns

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Appendix A. Creating External Commands in Linux

External commands allow you to add your own customized extensions to Mechanical APDL without relinking the program. You can create custom routines in C that access any of the Mechanical APDL API functions, link them into shared libraries using the supplied utilities, and execute the routines via the "external command" feature within Mechanical APDL. In addition, the program provides special commands that list all available external commands and allow you to reset all currently referenced external commands.

External command capability is supported on all Linux platforms. Refer to your ANSYS, Inc. Linux Installation Guide for currently supported compilers; the following instructions assume the presence of compatible compilers and linkers.

A.1. Tasks in Creating an External Command

To create a functional external command, you will need to complete the following general steps:

- · Create compilable source code.
- Create a shared library. This is facilitated by the gen_share utility and your system's make capability.
- Create an external table file (ans_ext.tbl), listing the various shared libraries, functions, and the related command.
- Set an environment variable pointing to the directory that holds the external table file.

The following sections detail each of these tasks.

A.1.1. Creating Compatible Code

You can create your functions using any of the API functions described in //ansys_inc/v211/ansys/customize/include/cAnsInterface.h, cAnsQuery.h, and cAnsPick.h. The following code segment demonstrates, at a minimal level, how to create functions that can be used as an entry point into a custom coded shared library.

The most important point in the following example it that the C program interface is an integer function that has one argument (a char pointer).

```
Parameters:
    Input
     uecmd
        The ANSYS external command string.
    Output
Return Value:
    The return value is ignored by the calling function;
       ----- End Function Description -----
int extfnc(char* uecmd)
   /* Note: uecmd is the entire command given to invoke this function */
   char* cmdsend = {"/COM, COMMAND SENT FROM EXTERNAL COMMAND"};
   char* querystr = {"NODE,,NUM,MAX"};
   char strrtn[32];
   int i, itype;
   double dblrtn;
   /* Send a simple command to be executed */
   i = cAnsSendCommand(cmdsend);
   /* Perform a simple query */
   i = cAnsGetValue(querystr,&dblrtn,strrtn,&itype);
   /* Display the value retrieved */
   cAns_printf("Max Node Number = %g\n",dblrtn);
   return (i);
```

A.1.2. Creating a Shared Library

Once you have written the source code for your functions, you can create a Makefile (using the **gen_share** utility) to build a shared library. The utility creates the Makefile in the current directory. The Makefile incorporates all the interdependencies of the C source files it encounters in that current directory. The **gen_share** utility is meant to setup the basic build. The user may need to make modifications to the Makefile depending on the situation.

The **gen_share** utility has the following syntax:

```
gen_share [-h] [-64] shared_object_name
```

where

-h

Produces command help.

-64

Configures the Makefile to use the -mips4 option for IRIX64.

shared_object_name

Is the name that will be given to the shared library.

As gen_share is executing, you may see one or more "No match" messages. This is normal. The script is searching for .c, .f, and .F file types in the current directory and returns this message if it cannot locate any files matching one of those types.

To create a shared library called mylibrary.so, you would issue the following command:

```
% gen_share mylibrary.so
```

The utility will produce a Makefile in the current directory. You will be able to generate the shared library by issuing the following command:

```
make
```

For example, to create the shared library for mylibrary.so, you would issue the following command:

```
% make
```

You will then find the specified shared library file in the current directory. You may also see warnings from the make process, and you may need to modify the Makefile or your source code.

A.1.3. Creating an External Table File

The external table file (ans_ext.tbl) can reside in any directory (but you must specify that directory in the ANSYS_EXTERNAL_PATH environment variable). The file contains an entry for each shared library function you wish to allow Mechanical APDL to access. There is no limit to the number of entries. The file entries have the following format:

```
/shared/library/path/library.so ~cm name function name
```

where:

/shared/library/path/library.so is the path to the directory that contains the shared library file. (Remotely mounted file systems are not recommended.)

~cm_name is the command used to invoke the function within Mechanical APDL. The command name must begin with a tilde (~) and each command name must be unique within the first four characters. The command name must be eight characters or less, including the tilde (~).

function_name is the name of the function that is referenced by the specified command name. (This must be unique within the first four characters if multiple external commands are specified.)

For example, the following entry references the /home/mydir/mylibs/myobject.so shared library and the myobject_function. It specifies ~myobj as the related command:

```
/home/mydir/mylibs/myobject.so ~myobj myobject_
```

Mechanical APDL also makes use of external commands, and places its own shared libraries and the associated external table file in the $/ansys_inc/v211/ansys/lib/<platform> directory (where <math><platform>$ is the directory specific to your computing platform, such as /linx64).

Mechanical APDL loads external commands as follows:

• Checks the ans_ext.tbl file in the /ansys_inc/v211/ansys/lib/<platform> directory and loads any external commands referenced there.

 Loads external commands referenced by the external table file in the directory designated with the ANSYS_EXTERNAL_PATH environment variable (see section Setting the ANSYS_EX-TERNAL_PATH Environment Variable (p. 366)).

If you designate a command name that has the same first four characters as a command listed in the $/ansys_inc/v211/ansys/lib/<platform>/ans_ext.tbl file, you will not be able to access your command. Therefore, it is a good practice to check the external table file to make sure you have no external command name conflicts. Do not modify the <math>/ansys_inc/v211/ansys/lib/<platform>/ans_ext.tbl file. You can also use the$ **~DEBUG**command to verify that no external command name conflicts exist.

Note:

The shared library must be consistent with the computer type and OS level on which Mechanical APDL is executed.

A.1.4. Setting the ANSYS_EXTERNAL_PATH Environment Variable

Before launching Mechanical APDL, first set the **ANSYS_EXTERNAL_PATH** to point to the directory containing the external table file. (For convenience, if you distribute your new functionality to other users they should set their .login or .cshrc files so that it is persistent from session to session.) For example, the following sets the environment variable to point to the /home/mydir directory.

setenv ANSYS_EXTERNAL_PATH /home/mydir

A.1.5. Using External Commands

To call an external command, enter it as you would any other Mechanical APDL command. You can also call external commands through either an APDL macro or UIDL script.

Note:

Avoid recursive external commands; that is, avoid situations where an external command calls another external command.

A.1.6. Checking External Command Status

You can check what shared libraries are currently accessible by entering the **~DEBUG** command in the command input window. The following figure shows an example of **~DEBUG** command output.

In this example, the output lists the command, the related shared library, the function, and if the command has been accessed.

A.1.7. Resetting External Commands

You can

- · Close all shared libraries
- · Free memory associated with external commands

by issuing the **~RESET** command. The command issues the following message to confirm that the reset operation was complete.

~RESET was processed: The external command buffers have been cleared.

Note:

The /CLEAR command also closes/resets all external command shared libraries.

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Appendix B. Creating External Commands in Windows

This section describes the steps required to create external commands on Windows platforms.

B.1. Tasks in Creating an External Command

To create a functional external command, you will need to complete the following general steps:

- Create compatible C source code.
- Create an external definition file (projname.def).
- Create a new project in Microsoft Visual Studio 2017.
- Create a shared library.
- Create an external table file (ans_ext.tbl), listing the various shared libraries, each function and the related command.
- Set the ANSYS_EXTERNAL_PATH environment variable

The following sections detail each of these tasks.

B.1.1. Creating Compatible Code

You can create your functions using any of the API functions described in Program Files\ANSYS Inc\V211\ansys\customize\include\cAnsInterface.h, cAnsQuery.h, and cAnspick.h. You can then execute these functions via the "external command" feature within Mechanical APDL. In addition, the program provides special commands that list all available external commands and allow you to reset all currently referenced external commands. The following code segment demonstrates, at a minimal level, how to create functions that can be used as an entry point into a custom coded shared library.

The most important point in the following example is:

The C program interface is an integer function that has one argument (a char pointer).

```
Parameters:
    Input
      uecmd
        The ANSYS external command string.
    Out.put.
Return Value:
    The return value is ignored by the calling function;
        ----- End Function Description -----
int extfnc(char* uecmd)
   /* Note: uecmd is the entire command given to invoke this function */
   char* cmdsend = {"/COM, COMMAND SENT FROM EXTERNAL COMMAND"};
   char* querystr = {"NODE,,NUM,MAX"};
       char strrtn[32];
   int i, itype;
   double dblrtn;
   /* Send a simple command to be executed */
   i = cAnsSendCommand(cmdsend);
   /* Perform a simple query */
   i = cAnsGetValue(querystr,&dblrtn,strrtn,&itype);
   /* Display the value retrieved */
   cAns_printf("Max Node Number = %g\n",dblrtn);
   return (i);
   }
```

B.1.2. Creating a Visual Studio Project

The steps for building a Visual Studio project are demonstrated in the example at the end of this appendix. See Example: Creating an External Command Using Visual Studio 2017 Professional (p. 373).

B.1.3. Creating an External Definition File

For each external function, you must declare it in the external definition file. The naming convention for this file is the name of your project with the .def extension; it must be located in your project directory. This file consists of the word *EXPORTS* on the first line, and the name(s) of the functions to be exported on each successive line. For the example function above:

```
EXPORTS extfunc
```

B.1.4. Creating a Shared Library

Once all of the necessary files have been incorporated into your project, simply compile (**Ctrl+F7**) and build (**F7**) the project. In your project directory, Developer Studio will create a Debug directory and will place the library in that directory (projname.dll).

B.1.5. Creating an External Table File

The external table file (ans_ext.tbl) can reside in any directory (but you must specify that directory in the ANSYS_EXTERNAL_PATH environment variable). The file contains an entry for each shared library function you wish Mechanical APDL to access. There is no limit to the number of entries. The file entries have the following format:

C:\shared\library\path\projname.dll ~cm_name function_name

where:

C:\shared\library\path\projname.dll is the path to the directory that contains the shared library file. (Remotely mounted file systems are not recommended.)

~cm_name is the command used to invoke the function within Mechanical APDL. The command name must begin with a tilde (~) and the first four characters of each command name must be unique.

function_name is the name of the function that is referenced by the specified command name. (This must be unique within the first four characters if multiple external commands are specified.)

For example, the following entry references the C:\home\mydir\mylibs\myobject.dll shared library and the myobject function, and specifies ~myobj as the related command:

```
C:\home\mydir\mylibs\myobject.dll ~myobj myobject
```

Mechanical APDL also makes use of external commands, and places its own shared libraries and the associated external table file in the C:\Program Files\ANSYS Inc\V211\ansys\lib\<platform> directory (where <platform> is the directory specific to your computing platform, such as \winx64). The program loads external commands in the following order:

- Checks the ans_ext.tbl file in the C:\Program Files\ANSYS Inc\V211\ansys\lib\<platform> directory and loads any external commands referenced there.
- Loads external commands referenced by the external table file in the directory designated with the ANSYS_EXTERNAL_PATH environment variable (see Setting the ANSYS_EXTERN-AL_PATH Environment Variable (p. 372)).

If you designate a command name that has the same first four characters as a command listed in the C:\Program Files\ANSYS Inc\V211\ansys\lib\<platform>\ans_ext.tbl file, you will not be able to access your command. Therefore, it is a good practice to check the external table file to make sure you have no external command name conflicts. Do not modify the C:\Program Files\ANSYS Inc\V211\ansys\lib\<platform>\ans_ext.tbl file. You can also use the ~DEBUG command to verify that no external command name conflicts exist.

Note:

The shared library must be consistent with the computer type and OS level on which Mechanical APDL is executed.

B.1.6. Setting the ANSYS_EXTERNAL_PATH Environment Variable

Before launching Mechanical APDL, you must first set the **ANSYS_EXTERNAL_PATH** to point to the directory containing the external table file. In Windows NT, the environment variables are in System Properties, which can be accessed through the Control Panel. For example, the following string sets the environment variable to point to the C:\home\mydir directory.

```
set ANSYS_EXTERNAL_PATH=C:\home\mydir
```

B.1.7. Using External Commands

To call an external command, issue it as you would any other Mechanical APDL command. You can also call external commands through either an APDL macro or UIDL routine.

Note:

Avoid recursive external commands; that is, avoid situations where an external command calls another external command.

B.1.8. Checking External Command Status

You can check what shared libraries are currently accessible by entering the **~DEBUG** command in the command input window. The following figure shows an example of **~DEBUG** command output.

Note that the output lists the command, the related shared library, the function, and whether or not the command has been accessed.

B.1.9. Resetting External Commands

You can

- · Close all shared libraries
- Free memory associated with external commands

by issuing the **~RESET** command. This command issues the following message to confirm that the reset operation is complete.

```
~RESET was processed: The external command buffers have been cleared.
```

Note:

The /CLEAR command also closes/resets all external command shared libraries.

B.1.10. Example: Creating an External Command Using Visual Studio 2017 Professional

An example for setting up an external command using Microsoft Visual Studio 2017 Professional is provided on the installation media. To run this example, perform the following steps.

- 1. Go to the Program Files\ANSYS Inc\V211\ANSYS\custom\user\winx64\ExtCmd directory.
- 2. Open the Visual Studio 2017 Professional solution file extcmd.sln (double click the file).
- 3. From the Visual Studio 2017 Professional menu, click on Build->Rebuild Solution.
- 4. Exit Visual Studio 2017 Professional.
- 5. Double-click on runextcmdtest.bat to run Mechanical APDL and test the external command that was just compiled.
- 6. In the output window enter ~excmd. You should see the following:

```
BEGIN:

-excmd

COMMAND SENT FROM EXTERNAL COMMAND

Max Node Number = 0

*** NOTE ***

CP = 0.625 TIME= 14:24:33

Command= -excmd was processed as an external command which is a non-standard use of the Mechanical APDL program.
```

7. Enter /exit, nosave to exit the program.

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Appendix C. User Material (UserMat) Subroutine Example

This example of a simple bilinear plasticity material model (identical to **TB**,BISO) demonstrates the user material subroutine UserMat, described in Subroutine UserMat (Creating Your Own Material Model) (p. 199).

C.1. UserMat Example Description

The example subroutine defines a 3-D material with plane strain and axisymmetric stress states. The analysis uses the 3-D solid element SOLID185. Comparison is made with the prediction by the **TB**,BISO material option.

The example is a two-element test case under simple tension. Element 1 has material defined using the **TB**,USER option, while Element 2 has material defined using the **TB**,BISO option. A 100-percent deformation is applied to both elements. Finite deformation (**NLGEOM**, ON) is considered. The POST26 processor results of stress components (Sxx, Syy) and plastic strain components (EPxx, EPyy) are printed for both elements. They are expected to be the same.

C.2. UserMat Example Input Data

```
/batch,list
/title, mvpl-um01, gal, usermat.F test case
/com, This is a single element test case for testing usermat.F
/com, usermat.F is the user materials subroutine
/com for current-technology elements.
/com, The material subroutine provided as the example
/com, is the same as the TB, BISO.
/com, A side by side comparison is made for two 185 elements,
/com, among which one is defined by TB,BISO, and another
/com, is defined as TB, USER. They are expected to produce
/com, the same results.
/com, uniaxial tension stress, large deformation.
/com,
/nopr
/nolist
/prep7
ele1=185
ele2=185
mat.1=1
mat2=2
et,1,ele1
keyopt, 1, 2, 1
mat.mat1
block,0,1,0,1,0,1
esize..1
vmesh,1
mat, mat2
block, 0, 1, 0, 1, 0, 1
```

```
esize,,1
vmesh,2
elist
! define material 1 by tb,biso
mp,ex , mat1,20e5
mp,nuxy,mat1,0.3
tb,biso,mat1,2,4
tbtemp,1.0
tbdata,1,1e3,100,
tbtemp,2.0
tbdata,1,2e3,100,
! define material 2 by tb,user
tb,user,mat2,2,4
tbtemp,1.0
                                    ! first temp.
tbdata,1,19e5, 0.3, 1e3,100,
                                    ! E, posn, sigy, H
tbtemp,2.0
tbdata,1,21e5, 0.3, 2e3,100,
tb, state, mat2,,8
                                    ! define 8 state variables
! boundary condition
nsel,s,loc,x
d,all,ux
nall
nsel,s,loc,y
d,all,uy
nall
nsel,s,loc,z
d,all,uz
nall
fini
/solu
tunif,1.5
nlgeom, on
nsel,s,loc,y,1
nsubst,20,100,1
d,all,uy,1.0
time,1
nall
outres,,-10
outpr,all,-10
solv
fini
/post26
eso1,2,1,,s,x,SX_BISO
esol,3,2,,s,x,SX_USER
esol,4,1,,s,y,SY_BISO
esol,5,2,,s,y,SY_USER
esol,6,1,,eppl,x,EPX_BISO
esol,7,2,,eppl,x,EPX_USER
esol,8,1,,eppl,y,EPY_BISO
esol,9,2,,eppl,y,EPY_USER
prvar, 2, 3, 4, 5
prvar,6,7,8,9
fini
/exit,no save
```

C.3. UserMat Example POST26 Output Results

```
**** ANSYS POST26 VARIABLE LISTING ****
               1 S
                            2 S X
                                          1 S Y
                                                        2 S
 TIME
                    Х
                                                             Y
               SX_BISO
                            SX_USER
                                          SY_BISO
                                                        SY_USER
             -0.188102E-02 -0.188102E-02 1509.45
0.10000
                                                       1509.45
0.28750
             -0.110968
                        -0.110968
                                          1525.07
                                                       1525.07
0.45625
             -0.814415
                          -0.814415
                                          1536.67
                                                       1536.67
0.66204
              -1.73160
                           -1.73160
                                          1548.95
                                                       1548.95
0.89592
              -1.86240
                            -1.86240
                                          1561.97
                                                        1561.97
1.0000
             -0.176924E-01 -0.176924E-01
                                         1569.16
                                                        1569.16
       ***** ANSYS POST26 VARIABLE LISTING *****
               1 EPPLX
                            2 EPPLX
                                          1 EPPLY
                                                        2 EPPLY
 TIME
              EPX_BISO
                            EPX_USER
                                          EPY_BISO
                                                       EPY_USER
0.10000
             -0.472687E-01 -0.472687E-01 0.945374E-01 0.945374E-01
             -0.125917
                        -0.125917
0.28750
                                                       0.251834
                                         0.251834
0.45625
             -0.187417
                          -0.187417
                                         0.374835
                                                       0.374835
0.66204
             -0.253409
                          -0.253409
                                         0.506818
                                                       0.506818
0.89592
             -0.319141
                          -0.319141
                                         0.638282
                                                       0.638282
             -0.345853
                          -0.345853
                                         0.691707
                                                       0.691707
1.0000
```

C.4. USERMAT.F List File for This Example

```
subroutine usermat(
                          matId, elemId, kDomIntPt, kLayer, kSectPt,
     &
                         ldstep, isubst, keycut,
                         nDirect, nShear, ncomp, nStatev, nProp,
     ۶
                         Time, dTime, Temp, dTemp,
                         stress, ustatev, dsdePl, sedEl, sedPl, epseq,
                         Strain, dStrain, epsPl, prop, coords,
                         var0, defGrad_t, defGrad,
                         tsstif, epsZZ,
                         var1, var2, var3, var4, var5,
                         var6, var7, var8)
C*******
      *** primary function ***
C
С
            user defined material constitutive model
С
С
      Attention:
С
            User must define material constitutive law properly
C
            according to the stress state such as 3D, plane strain
С
            and axisymmetry, plane stress and 3D/1D beam.
С
C
            a 3D material constitutive model can use for
С
С
            plane strain and axisymmetry cases.
C
            When using shell elements, a plane stress algorithm
С
            must be use.
C
С
С
        The following demonstrates a USERMAT subroutine for
C
        a plasticity model, which is the same as TB, BISO,
С
        for different stress states.
С
С
        See "ANSYS user material subroutine USERMAT" for detailed
        description of how to write a USERMAT routine.
С
С
        This routine calls four routines,
С
        usermat3d.F, usermatps.F usermatbm.F and usermat1d.F, w.r.t.
C
С
        the corresponding stress states.
        Each routine can be also a usermat routine for the specific
```

```
С
        element.
С
C*
С
С
      input arguments
C
      ==========
      matId
С
                (int,sc,i)
                                         material #
С
       elemId
                (int,sc,i)
                                         element #
                                         "k"th domain integration point
       kDomIntPt (int,sc,i)
C
С
      kLayer
                (int,sc,i)
                                         "k"th layer
                                         "k"th Section point
      kSectPt
                (int,sc,i)
С
С
       ldstep
                (int,sc,i)
                                         load step number
С
       isubst
                (int,sc,i)
                                         substep number
      nDirect
                (int,sc,in)
                                         # of direct components
С
С
      nShear
                (int,sc,in)
                                         # of shear components
С
      ncomp
                (int,sc,in)
                                         nDirect + nShear
C
      nstatev
                (int,sc,1)
                                         Number of state variables
                                         Number of material ocnstants
С
       nProp
                (int,sc,1)
С
      Temp
                                         temperature at beginning of
С
                (dp,sc,in)
                                         time increment
С
       dTemp
                                         temperature increment
С
                 (dp,sc,in)
       Time
                (dp,sc,in)
                                         time at beginning of increment (t)
С
       dTime
                (dp,sc,in)
                                         current time increment (dt)
С
C
С
       Strain
               (dp,ar(ncomp),i)
                                         Strain at beginning of time increment
С
       dStrain (dp,ar(ncomp),i)
                                         Strain increment
       prop
                (dp,ar(nprop),i)
                                         Material constants defined by TB, USER
C
С
       coords
                (dp,ar(3),i)
                                         current coordinates
       defGrad_t(dp,ar(3,3),i)
С
                                         Deformation gradient at time t
      defGrad (dp,ar(3,3),i)
                                         Deformation gradient at time t+dt
C
С
С
      input output arguments
      C
       stress (dp,ar(nTesn),io)
С
                                         stress
                                         user state variables
               (dp,ar(nstatev),io)
С
       ustatev
С
       sedEl
               (dp,sc,io)
                                         elastic work
       sedPl
                (dp,sc,io)
                                         plastic work
С
                                         equivalent plastic strain
       epseq
                (dp,sc,io)
C
С
       tsstif
                (dp,ar(2),io)
                                         transverse shear stiffness
                                         tsstif(1) - Gxz
С
                                         tsstif(2) - Gyz
С
                                         tsstif(1) is also used to calculate hourglass
С
С
                                         stiffness, this value must be defined when low
С
                                         order element, such as 181, 182, 185 with uniform
                                         integration is used.
С
                                         not used, they are reserved arguments
                (dp,sc,io)
C
       var?
С
                                         for further development
C
      output arguments
C
С
С
       keycut (int,sc,io)
                                         loading bisect/cut control
                                         0 - no bisect/cut
C
                                         1 - bisect/cut
С
                                         (factor will be determined by ANSYS solution control)
С
С
       dsdePl
                (dp,ar(ncomp,ncomp),io)
                                         material jacobian matrix
       epsZZ
                (dp,sc,o)
                                         strain epsZZ for plane stress,
С
                                         define it when accounting for thickness change
С
                                         in shell and plane stress states
С
С
   *******************
C*
С
С
       ncomp
                  for 3D (nshear=3)
                  for plane strain or axisymmetric (nShear = 1)
C
       ncomp
                  for plane stress (nShear = 1)
С
              3
                  for 3d beam
                                 (nShear = 2)
С
       ncomp
С
              1
                  for 1D (nShear = 0)
С
       stresss and strains, plastic strain vectors
С
          11, 22, 33, 12, 23, 13 for 3D
С
          11, 22, 33, 12
                                    for plane strain or axisymmetry
```

```
11, 22, 12
                                  for plane stress
С
                                  for 3d beam
С
          11, 13, 12
                                  for 1D
С
          11
С
С
      material jacobian matrix
        3D
C
           dsdePl
                      1111
                             1122
                                   1133
                                          1112
                                                1123
С
                            2222
                                                      2213
           dsdePl
                      2211
                                   2233
                                          2212
                                                2223
С
           dsdePl
                       3311
                             3322
                                   3333
                                         3312
                                                3323
                                                      3313
C
С
           dsdePl
                      1211
                             1222
                                   1233
                                         1212
                                                1223
                                                      1213
                    2311
                             2322
                                   2333
                                          2312
           dsdePl
                                                2323
                                                      2313
С
                             1322
                                   1333
                                          1312
С
           dsdePl
                                                1323
                                                       1313
        plane strain or axisymmetric (11, 22, 33, 12)
С
                    1111 1122
                                   1133 1112
          dsdePl
С
           dsdePl
                    2211
                             2222
                                    2233
                                         2212
С
                    3311
           dsdePl
                           3322
                                   3333
                                         3312
С
           dsdePl
                     1211
                             1222
                                   1233
                                          1212
С
        plane stress (11, 22, 12)
С
                    | 1111
С
          dsdePl
                             1122
                                    1112
                    2211
          dsdePl
                             2222
                                    2212
С
           dsdePl
                    1211
                             1222
                                   1212
С
        3d beam (11, 13, 12)
С
                  1111
С
           dsdePl
                             1113
                                    1112
С
           dsdePl
                       1311
                             1313
                                    1312
                    1211
                             1213
           dsdePl
                                   1212
C
С
С
           dsdePl
                    | 1111 |
C
#include "impcom.inc"
C
     INTEGER
                     matId, elemId,
    &
                     kDomIntPt, kLayer, kSectPt,
    &
                     ldstep, isubst, keycut,
    &
                     nDirect,nShear,ncomp,nStatev,nProp
     DOUBLE PRECISION
                     Time,
                             dTime, Temp,
                                              dTemp,
                     sedEl,
                             sedPl, epseq,
                                            epsZZ
     DOUBLE PRECISION
                     stress (ncomp ), ustatev (nStatev),
    &
                     dsdePl (ncomp, ncomp),
    &
                     Strain (ncomp ), dStrain (ncomp ),
    δ
                     epsPl (ncomp ), prop (nProp ),
    &
                     coords (3),
                     defGrad_t(3,3),
                                      defGrad(3,3),
                     tsstif (2)
С
c********** User defined part *******************************
C
c --- parameters
С
     INTEGER
                    NEWTON, mcomp
     DOUBLE PRECISION HALF, THIRD, ONE, TWO, SMALL, ONEHALF,
                     ZERO, TWOTHIRD, ONEDM02, ONEDM05, sqTiny
    æ
                               = 0.d0,
     PARAMETER
                    (ZERO
                               = 0.5d0,
    &
                     HALF
                               = 1.d0/3.d0,
                     THIRD
    &
                               = 1.d0,
                     ONE
    &
    &
                     TWO
                               = 2.d0,
                     SMALL
                               = 1.d-08,
    &
                     sqTiny
                               = 1.d-20,
                     ONEDM02
                               = 1.d-02,
    &
                     ONEDM05
                               = 1.d-05,
                     ONEHALF
                               = 1.5d0,
                     TWOTHIRD
                               = 2.0d0/3.0d0,
                     NEWTON
                               = 10,
                     mcomp
                               = 6
c --- local variables
```

```
C
                (dp,ar(6),1)
                                         trial stress
С
       sigElp
       dsdeEl
                (dp,ar(6,6),1)
                                         elastic moduli
С
       sigDev
                (dp,ar(6),1)
                                         deviatoric stress tensor
С
С
       dfds
                (dp,ar(6),1)
                                         derivative of the yield function
       JM
                (dp,ar(6,6),1)
                                         2D matrix for a 4 order tensor
C
       pEl
                (dp,sc
                                         hydrostatic pressure stress
С
                          ,1)
С
       qEl
                (dp,sc
                          ,1)
                                         von-mises stress
                          ,1)
                                         equivalent plastic strain at
С
       pleq_t
                (dp,sc
С
                                         beginnig of time increment
               (dp,sc
                                         equivalent plastic strain at end
С
       pleq
                          ,1)
С
                                         of time increment
С
       dpleq
                (dp,sc
                          ,1)
                                         incremental equivalent plastic strain
                                         correction of incremental
                (dp,sc
С
       cpleq
                          ,1)
С
                                         equivalent plastic strain
С
       sigy_t
               (dp,sc
                          ,1)
                                         yield stress at beginnig of time increments
                          ,1)
                                         yield stress at end of time
       sigy
               (dp,sc
С
                                         increment
С
С
       young
                (dp,sc
                          ,1)
                                         Young's modulus
                          ,1)
                (dp,sc
                                         Poiss's ratio
C
       posn
       sigy0
                (dp,sc
                          ,1)
                                         initial yield stress
С
                          ,1)
                                         plastic slope
       dsigdep
               (dp,sc
С
       twoG
                (dp,sc
                          ,1)
                                         two time of shear moduli
С
       threeG
                (dp,sc
                          ,1)
                                         three time of shear moduli
С
                                         nonlinear function to be solved
C
       funcf
                (dp,sc
                          ,1)
С
                                         for dpleq
С
       dFdep
                (dp,sc
                          ,1)
                                         derivative of nonlinear function
С
                                         over dpleq
С
c --- temporary variables for solution purpose
C
       threeOv2qEl, oneOv3G, qElOv3G, con1, con2, fratio
С
C
      {\tt DOUBLE\ PRECISION\ sigElp(mcomp),\ dsdeEl(mcomp,mcomp),\ G(mcomp),}
                      sigDev(mcomp), JM (mcomp,mcomp), dfds(mcomp)
      DOUBLE PRECISION var0, var1, var2, var3, var4, var5,
                      var6, var7, var8
      DATA G/1.0D0,1.0D0,1.0D0,0.0D0,0.0D0,0.0D0/
С
      INTEGER
                      i, j
                            qEl,
     DOUBLE PRECISION pEl,
                                      pleq_t, sigy_t , sigy,
                      cpleq, dpleq,
                                      pleq,
                      young, posn,
                                      sigy0,
                                               dsigdep,
     &
     &
                      elast1, elast2,
                      twoG, threeG, oneOv3G, qElOv3G, threeOv2qEl,
     δ
                      funcf, dFdep, fratio, con1, con2
     δ
C
      keycut = 0
      dsigdep = ZERO
      pleq_t = ustatev(1)
             = pleq_t
     pleq
c *** get Young's modulus and Poisson's ratio, initial yield stress and others
      young = prop(1)
              = prop(2)
      posn
      sigy0
              = prop(3)
c *** calculate the plastic slope
      dsigdep = young*prop(4)/(young-prop(4))
              = young / (ONE+posn)
      twoG
      threeG
             = ONEHALF * twoG
С
c *** calculate elastic stiffness matrix (3-D)
С
С
      elast1=young*posn/((1.0D0+posn)*(1.0D0-TW0*posn))
      elast2=young/(TWO*(1.0D0+posn))
      dsdeEl(1,1) = (elast1+TWO*elast2)*G(1)*G(1)
      dsdeEl(1,2)=elast1*G(1)*G(2)+elast2*TWO*G(4)*G(4)
      dsdeEl(1,3)=elast1*G(1)*G(3)+elast2*TWO*G(5)*G(5)
```

```
dsdeEl(1,4)=elast1*G(1)*G(4)+elast2*TWO*G(1)*G(4)
      dsdeEl(1,5) = elast1*G(1)*G(5) + elast2*TWO*G(1)*G(5)
      dsdeEl(1,6)=elast1*G(1)*G(6)+elast2*TWO*G(4)*G(5)
      dsdeEl(2,2) = (elast1+TWO*elast2)*G(2)*G(2)
      dsdeEl(2,3)=elast1*G(2)*G(3)+elast2*TWO*G(6)*G(6)
      dsdeEl(2,4)=elast1*G(2)*G(4)+elast2*TWO*G(1)*G(4)
      dsdeEl(2,5)=elast1*G(2)*G(5)+elast2*TWO*G(1)*G(5)
      dsdeEl(2,6)=elast1*G(2)*G(6)+elast2*TWO*G(2)*G(6)
      dsdeEl(3,3) = (elast1+TWO*elast2)*G(3)*G(3)
      dsdeEl(3,4)=elast1*G(3)*G(4)+elast2*TWO*G(5)*G(6)
      dsdeEl(3,5)=elast1*G(3)*G(5)+elast2*TWO*G(5)*G(3)
      dsdeEl(3,6) = elast1*G(3)*G(6) + elast2*TWO*G(6)*G(3)
      dsdeE1(4,4)=elast1*G(4)*G(4)+elast2*(G(1)*G(2)+G(4)*G(4))
      dsdeEl(4,5) = elast1*G(4)*G(5) + elast2*(G(1)*G(6)+G(5)*G(4))
      dsdeEl(4,6)=elast1*G(4)*G(6)+elast2*(G(4)*G(6)+G(5)*G(2))
      dsdeEl(5,5)=elast1*G(5)*G(5)+elast2*(G(1)*G(3)+G(5)*G(5))
      dsdeE1(5,6)=elast1*G(5)*G(6)+elast2*(G(4)*G(3)+G(5)*G(6))
      dsdeEl(6,6)=elast1*G(6)*G(6)+elast2*(G(2)*G(3)+G(6)*G(6))
      do i=1,ncomp-1
        do j=i+1,ncomp
          dsdeEl(j,i)=dsdeEl(i,j)
        end do
      end do
С
c *** calculate the trial stress and
      copy elastic moduli dsdeEl to material Jacobian matrix
С
      do i=1,ncomp
         sigElp(i) = stress(i)
         do j=1,ncomp
            dsdePl(j,i) = dsdeEl(j,i)
            sigElp(i) = sigElp(i)+dsdeEl(j,i)*dStrain(j)
         end do
      end do
c *** hydrostatic pressure stress
      pEl = -THIRD * (sigElp(1) + sigElp(2) + sigElp(3))
c *** compute the deviatoric stress tensor
      sigDev(1) = sigElp(1) + pEl
      sigDev(2) = sigElp(2) + pEl
      sigDev(3) = sigElp(3) + pEl
      sigDev(4) = sigElp(4)
      sigDev(5) = sigElp(5)
      sigDev(6) = sigElp(6)
c *** compute von-mises stress
      qEl =
     & sigDev(1) * sigDev(1)+sigDev(2) * sigDev(2)+
     & sigDev(3) * sigDev(3)+
     & TWO*(sigDev(4) * sigDev(4)+ sigDev(5) * sigDev(5)+
     & sigDev(6) * sigDev(6))
      qEl = sqrt( ONEHALF * qEl)
c *** compute current yield stress
              = sigy0 + dsigdep * pleq
      sigy
С
      fratio = qEl / sigy - ONE
c *** check for yielding
      IF (sigy .LE. ZERO.or.fratio .LE. -SMALL) GO TO 500
C
      sigy_t = sigy
      threeOv2qEl = ONEHALF / qEl
c *** compute derivative of the yield function
      DO i=1, ncomp
        dfds(i) = threeOv2qEl * sigDev(i)
      END DO
      oneOv3G = ONE / threeG
      qElOv3G = qEl * oneOv3G
c *** initial guess of incremental equivalent plastic strain
               = (qEl - sigy) * oneOv3G
      dpleq
      pleq
               = pleq_t + dpleq
c *** Newton-Raphson procedure for return mapping iteration
      DO i = 1, NEWTON
         sigy = sigy0 + dsigdep * pleq
```

```
funcf = qElOv3G - dpleq - sigy * oneOv3G
         dFdep = - ONE - dsigdep * oneOv3G
         cpleq = -funcf / dFdep
        dpleq = dpleq + cpleq
С
         --- avoid negative equivalent plastic strain
        dpleq = max (dpleq, sqTiny)
              = pleq_t + dpleq
        pleq
         fratio = funcf/qElOv3G
        check covergence
        IF (((abs(fratio) .LT. ONEDM05
              (abs(cpleq ) .LT. ONEDM02 * dpleq)) .OR.
    æ
    &
             ((abs(fratio) .LT. ONEDM05
                                         ) .AND.
              (abs(dpleq ) .LE. sqTiny
                                              ))) GO TO 100
     END DO
c *** Uncovergence, set keycut to 1 for bisect/cut
     keycut
      GO TO 990
100 CONTINUE
c *** update stresses
      con1 = twoG * dpleq
     DO i = 1 , ncomp
         stress(i) = sigElp(i) - con1 * dfds(i)
     END DO
С
c *** update plastic strains
     DO i = 1 , nDirect
        epsPl(i) = epsPl(i) + dfds(i) * dpleq
     END DO
     DO i = nDirect + 1 , ncomp
        epsPl(i) = epsPl(i) + TWO * dfds(i) * dpleq
      END DO
     epseq = pleq
c *** Update state variables
     ustatev(1) = pleq
c *** Update plastic work
     sedPl = sedPl + HALF * (sigy_t+sigy)*dpleq
C
c *** Material Jcobian matrix
С
      IF (qEl.LT.sqTiny) THEN
        con1 = ZERO
      ELSE
        con1 = threeG * dpleq / qEl
      END IF
     con2 = threeG/(threeG+dsigdep) - con1
     con2 = TWOTHIRD * con2
     DO i=1, ncomp
        DO j=1,ncomp
           JM(j,i) = ZERO
        END DO
      END DO
      DO i=1,nDirect
        DO j=1,nDirect
           JM(i,j) = -THIRD
        END DO
        JM(i,i) = JM(i,i) + ONE
      END DO
      DO i=nDirect + 1,ncomp
        JM(i,i) = HALF
      END DO
     DO i=1,ncomp
        DO j=1,ncomp
            dsdePl(i,j) =
                           dsdeEl(i,j) - twoG
                 * ( con2 * dfds(i) * dfds(j) + con1 * JM(i,j) )
        END DO
     END DO
С
     goto 600
  500 continue
```

```
c *** Update stress in case of elastic/unloading
    do i=1,ncomp
        stress(i) = sigElp(i)
    end do

600 continue
c *** Claculate elastic work
    sedEl = ZERO
    DO i = 1 , ncomp
        sedEl = sedEl + stress(i)*(Strain(i)+dStrain(i)-epsPl(i))
    END DO
    sedEl = sedEl * HALF

c
    return
    end
```

C.5. Accessing Solution and Material Data

These APIs are provided for your convenience to help you access solution and material data easily.

```
c *** subroutine get_ElmInfo(inquire, value)
С
         description
С
С
            function to inquire element and solution information
         definition
C
С
            inquire - query argument (string)
С
            value - value of query argument
         variables
C
С
           inquire
                           - value
С
            'LDSTEP'
                           - load step number
            'ISUBST'
                           - substep step number
C
            'IEQITR'
                           - current interation number
С
С
            'NUMINTG'
                           - number of gauss integration
            'ELEMID'
                           - element number
C
С
            'MATID'
                           - material number of current element
                           - number of state variable for current material at
            'NSVAR'
C
С
                             gauss intg.
С
            'NCOMP'
                           - number of vector components, such as stresses
С
c *** subroutine get_ElmData (kchar, elemId, kMatRecPt, ncomp, vect)
C
С
         description
            function to get/inquire solution dependent variables
С
            such as stress, strains at gauss intg. point.
С
         definition
С
           kchar
C
                           - string variable containing a query argument
С
            elemId
                           - element number
            kMatRecPt
                           - material integration point number
С
                           - number of components to be inquired
C
            ncomp
С
                               Use the 'NCOMP' query to determine the correct
С
                               array sizes for tensor quantities
                           - variable array containing the retrieved variables
           vect(*)
C
С
         variables
                           - element coordinate systems
С
           'ESYS'
С
           'ISIG'
                           - initial stress
           'SIG'
                           - stress (for CPT elements with pore pressure DOFs, this query returns effective stress)
С
           'EPTO'
                           - total strain
С
                           - plastic strain
           'EPPL'
С
С
           'EPCR'
                           - creep strain
           'EPTH'
                           - thermal strain
C
                           - swelling strain
С
           'EPSW'
С
           'SVAR'
                           - state variables
           'PLEO'
                           - accumulated equivalent plastic strain
C
```

```
- accumulated equivalent creep strain
С
                           - total stress (this is available only for CPT elements with pore pressure)
С
           'TSIG'
С
c *** subroutine put_ElmData (inquire, elemId, kIntg, nvect, vect)
        description
С
           function to put solution dependent variables
С
            such as stress, strains at gauss intg. point.
С
         !! Use this in caution, it overides ansys database. Usually
C
С
         !! you should only write user defined state variables,
         !! SVAR
С
         definition
С
С
            inquire
                           - query argument (string)
                           - element number
            elemId
С
           kIntg
                           - gauss intg. number
С
С
           nvect
                           - number of vector to be inquired
                               Use the 'NCOMP' query to determine the correct
С
                               array sizes for tensor quantities
С
С
            vect
                           - vector to be inquired
С
С
         variables
           'SIG '
                           - stress vector
С
С
           'EPTO'
                           - Total strain vector (EPEL+EPPL+EPCR+EPTH)
С
           'EPPL'
                           - plastic strain vector
           'EPCR'
                           - creep strain vector
С
                           - thermal strain vector
С
           'EPTH'
           'ISIG'
С
                           - Initial stress vector
С
           'PLEQ'
                           - accumulated equivalent plastic strain
                           - accumulated equivalent creep strain
           'CREQ'
С
           'SVAR'
                           - State variables (define by tb, state)
С
```

Appendix D. Structural-Thermal User Material (UserMat, UserMatTh) Example

This example demonstrates user-defined structural and thermal materials in a coupled structural-thermal analysis.

D.1. Example Description

The problem considered in the example is a 3-D version of the verification test case VM296 in the ANSYS Mechanical APDL Verification Manual.

The bilinear isotropic hardening plastic material with temperature-dependent yield stress and tangent modulus is defined using the **TB**,USER model. The linear thermal material is defined using **TB**,USER,,,THERM. The source files for the UserMat.F and UserMatTH.F subroutines used in this example can be found in subdirectory: /ansys_inc/v195/ansys/customize/user/ (Linux) or Program Files\ANSYS Inc\V195\ansys\customize\user\ (Windows).

The model is discretized using SOLID226 elements.

Results are expected to agree closely with the results of a 2-D simulation (VM296) performed for simulation time = 13 s.

D.2. Example Input Data

```
/TITLE, USER MATERIAL FOR RADIAL EXPANSION OF A THERMOPLASTIC CYLINDER
/COM, REFERENCE: "THERMOMECHANICAL MODELING OF METALS AT FINITE STRAINS:
/COM, FIRST AND MIXED ORDER FINITE ELEMENTS", LAURENT ADAM, JEAN-PHILIPPE PONTHOT,
/COM, INTERNATIONAL JOURNAL OF SOLIDS AND STRUCTURES 42 (2005) 5615-5655
/COM.
RI=0.1
                     ! INTERNAL RADIUS
RE=0.2
                     ! EXTERNAL RADIUS
DX = 0.13
                     ! RADIAL DISPLACEMENT
TREF=293
                     ! REFERENCE TEMPERATURE, K
/UPF,usermat.F
/UPF, usermatth.F
/PREP7
ET.1.SOLID226.11
                         ! STRUCTRAL-THERMAL COUPLING
KEYOT, 1, 9, 1
                          ! SUPPRESSED THERMOELASTIC DAMPING
MP,ALPX,1,23.8E-6
                          ! THERMAL EXPANSION COEFFICIENT
MP, DENS, 1, 2700
                          ! DENSITY
MP,QRATE,1,0.9
                          ! TAYLOR-QUINNEY FACTOR
TB, USER, 1, 6, 4
                          ! USER-DEFINED STRUCTURAL MATERIAL
TBTEMP, 293
TBDATA,1,7.E10,0.3,70E6,210E6
                                   ! E, NU, SIGY, ET
TBTEMP,313
TBDATA,1,7.E10,0.3,69.58E6,210E6 ! E,NU,SIGY,ET
```

```
TBTEMP, 333
TBDATA,1,7.E10,0.3,69.16E6,210E6 ! E,NU,SIGY,ET
TBTEMP.353
TBDATA,1,7.E10,0.3,68.74E6,210E6 ! E,NU,SIGY,ET
TBTEMP, 373
TBDATA, 1, 7. E10, 0.3, 68.32E6, 210E6 ! E, NU, SIGY, ET
TBTEMP,393
TBDATA,1,7.E10,0.3,67.9E6,210E6 ! E,NU,SIGY,ET
TB, USER, 1, 1, 2, THERM
                          ! USER-DEFINED THERMAL MATERIAL
TBDATA,1,150,900
                          ! K,C
TB, STATE, 1,,8
                          ! EPEQ, EPPL, SEQV
RECT, RI, RE, 0, RE-RI
TYPE,1
MAT,1
LESIZE, 1,,,10
LESIZE, 2,,,1
LESIZE, 3,,,10
LESIZE, 4,,,1
K,100,0
K,101,0,1
VROT,1,,,,,,100,101,10,1
VMESH,1
ALLSEL, ALL
CSYS,5
NROTATE, ALL
NSEL,S,LOC,z,0
NSEL, A, LOC, z, RE-RI
D,ALL,Uz,0
                         ! CONSTRAIN UY DOF AT TOP AND BOTTOM FACES TO
                         ! SIMULATE INFINITELY LONG CYLINDER
NSEL,S,LOC,X,RI
D,ALL,UX,DX ! APPLY RADIAL DISPLACEMENT ALONG INNER SURFACE OF CYLINDER
NSEL, S, LOC, Y, 0
NSEL, A, LOC, Y, 10
D, ALL, UY
ALLSEL, ALL
CSYS,0
TREF, TREF ! REFERENCE TEMPERATURE
IC, ALL, TEMP, TREF
                   ! INITIAL CONDITION
FINISH
/COM.
/COM, **** SOLUTION TIME = 13 SEC (ADIABATIC PROCESS) *****
/COM,
/SOLUTION
ANTYPE, TRANSIENT
                       ! TRANSIENT ANALYSIS
NLGEOM, ON
                       ! LARGE DEFLECTION ON
TINTP,,,,1.0
                       ! TRANSIENT INTEGRATION PARAMETER
KBC,0
                       ! RAMPED LOADING
                       ! NUMBER OF SUBSTEPS
NSUB, 20, 100, 5
OUTRES, ALL, ALL
CNVTOL, HEAT, 1.E-2, 1.E-2 ! CONVERGENCE TOLERANCE
TIME, 13 ! END TIME
SOLVE
FINISH
/POST26
NSEL,S,LOC,X,0.1
NSEL, R, LOC, Y, 0.1
ND=NDNEXT(0)
NSOL, 2, ND, TEMP
                        ! TEMPERATURE AT NODE =ND
FILLDATA, 3, , , , -1
FILLDATA, 4, , , , 293.0
PROD, 5, 3, 4
ADD,6,2,5
NSOL, 7, ND, U, X, UX ! UX AT NODE=ND
```

```
PRVAR,7,6
/GRID,1
/AXLAB,X,Displacement of inner surface[m] /AXLAB,Y,Temperature variation [K]
/XRANGE,0,0.14
/GROPT,DIVX,14
/YRANGE,0,60
/GROPT,DIVY,6
XVAR,7
PLVAR,6
ALLSEL, ALL
FINISH
/POST1
SET, LAST, LAST
PLNSOL, TEMP
PLNSOL, EPPL, EQV
FINISH
```

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Appendix E. Fully Coupled Wind Turbine Example in Mechanical APDL

This wind coupling example solution has been implemented in Mechanical APDL to illustrate how to perform an integrated analysis of a wind turbine with its supporting structure. In this solution procedure, both the structural code (that is, Mechanical APDL) and aeroelastic (e.g. Flex5) code are run simultaneously with continuous data exchange between the two programs at each time step. The data transfer is done through a set of interface routines that put or get data from a shared common data space. The interface is supplied in the form of a DLL which both the structural and aeroelastic programs will be accessing during an analysis. The example includes user programmable functions, and a macro has been developed to facilitate the communication with the DLL from Mechanical APDL. Aeroelastic code developers will need to utilize the same set of routines to establish communications between their code and the DLL.

There is also a sequential coupling method available; see Sequential Coupled Wind Turbine Solution in Mechanical APDL in the *Advanced Analysis Guide* for more information about using that method.

E.1. Implementing a Fully Coupled Wind Turbine Analysis

This example implementation of the fully coupled wind solution in Mechanical APDL follows a similar strategy to that used for ASAS. In particular, data access is provided through the same set of interface routines. This enables easy adaptation of the new facility by existing ASAS wind turbine users.

The following summarize the modeling characteristics in Mechanical APDL for a wind coupling analysis:

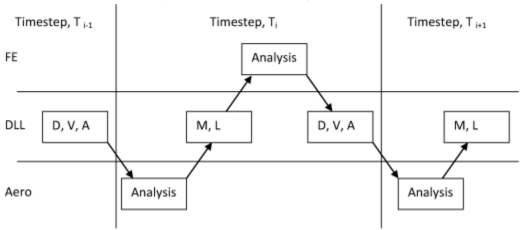
- The turbine effect is modeled via the user element USER300. This special user element has 9 nodes, with 6 freedoms (UX, UY, UZ, ROTX, ROTY, and ROTZ) on the first node and 3 freedoms (UX, UY, and UZ) on each of the subsequent nodes, making it capable of having a maximum 30 degrees of freedom on the element. The first node is the connection point between the turbine and the supporting structure and therefore it must be a node in the structural model. The other 8 nodes are created to accommodate the additional freedoms that are internal to the turbine element and are used solely by the aeroelastic code. Therefore, these nodes should not be connected to any other parts of the model.
- Key option 1 (KEYOPT(1)) of the user element is used to specify the damping matrix option. Damping can be obtained from the aeroelastic code alone, or computed from Rayleigh damping in Mechanical APDL based on the turbine mass and stiffness matrices, or both. KEYOPT(1) = 0 indicates that the damping matrix will be taken from the aeroelastic code plus Rayleigh damping, and this is the default. KEYOPT(1) = 1 indicates that only Rayleigh damping will be used. KEYOPT(1) = 2 indicates that only damping from the aeroelastic code will be used.
- The element does not have any material property or real constant data.
- The element mass will not generate any body forces even if accelerations (e.g. ACEL) are defined.

- The only element results available are the element nodal forces.
- A Mechanical APDL command macro called **WTBCREATE** is provided to assist with the creation of a wind turbine model. This will automatically generate a turbine element and issue relevant data commands that are necessary to run a wind coupling analysis.
- Special versions of the user subroutines UserElem, USolBeg, USsFin and USolFin are provided to enable a wind coupling analysis. In addition, the shared common DLL WTBFunctions.dll is also required. A custom build of Mechanical APDL is required during which the aeroelastic linking option should be selected.
- The analysis type should be transient (ANTYPE, TRANS) for a wind coupling analysis.

E.2. Theory

The whole offshore wind turbine structure is split into two parts. The upper part is the rotor-nacelle-assembly of the turbine including the tower and this is modeled by a wind turbine aeroelastic code. The lower part is the support structure and this is modeled by structural FE as usual.

In this coupling approach, both the aeroelastic and FE code runs concurrently, with data exchange between the two programs occurring at every time step. The turbine effect as modeled by the aeroelastic code is taken into account in the FE code as a special element, which accepts the wind turbine system matrices (stiffness, damping and mass) and aerodynamic load vector as if it is a superelement.



In the diagram above, the following data items are exchanged:

D, V, A: Displacements, Velocities, Acceleration information at interface point.

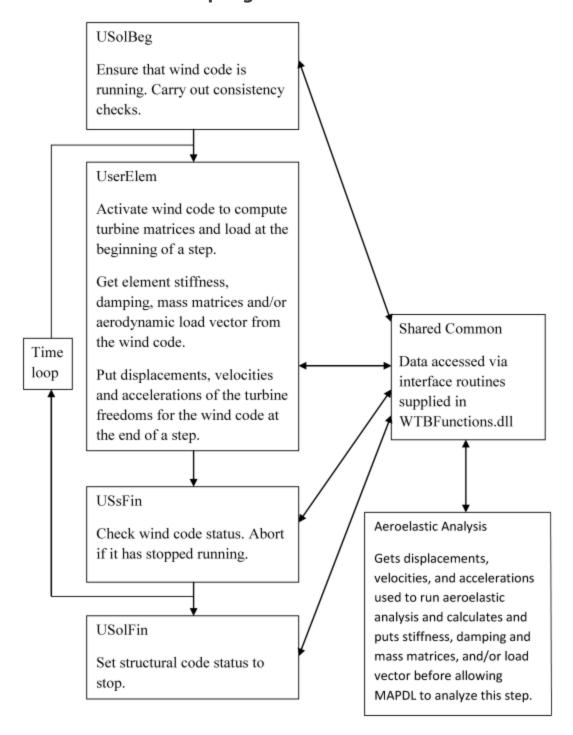
M, L: Matricies (Stiffness, Damping and Mass), and/or Load Vectors.

At the beginning of a time step, the aeroelastic code is called to compute the turbine matrices and loading based on the kinematics at the end of the previous step. It is assumed that the turbine data will remain constant during the time step. After transferring the information to the FE code, the full system of equations of motion is solved by the FE code to obtain an updated solution. Finally, the turbine kinematics are passed back to the aeroelastic code to continue the time advancement process.

In the case of Flex5, the maximum number of equations generated by a model is 28.

Further details about the theory of the fully coupled solution can be obtained from the work of Kaufer [1] and Seidal [2].

Flowchart of Wind Coupling Calculations



E.3. Compiling a Custom Version of Mechanical APDL

To activate the wind coupled analysis feature, it is necessary to create a custom build of Mechanical APDL that includes the wind interface libraries Aeroelastic.lib and WTBFunctions.lib.The

library Aeroelastic.lib contains updated user routines to enable a wind coupling analysis in Mechanical APDL. The library WTBFunctions.lib contains interface routines for storing and retrieving data to a shared common area. Since the wind coupling solution involves changes to some of the user routines, it is important that the library Aeroelastic.lib is linked in before all other user libraries so that the wind library versions are picked up by the linker.

The executable of the aeroelastic code should be placed in the same folder as the custom Mechanical APDL executable as both programs need to use WTBFunctions.dll to access the shared data. Alternatively, the folder containing the WTBFunctions.dll could be added to the path (on Windows systems) to enable the aeroelastic code to remain elsewhere.

To link a custom version, run ANS_ADMIN from the utilities folder in the start menu. During the relinking, the user is asked if the aeroelastic functions are to be included.

For further information and compiler requirements regarding the linking procedure and additional settings for using custom compiled executables, see Compiling and Linking UPFs on Windows Systems in the *Programmer's Reference* (p. 138).

E.4. Performing a Wind Coupled Analysis

In order to perform a wind coupling analysis, the aeroelastic software must be modified to provide data communication with Mechanical APDL through the specified interface.

The aeroelastic analysis should be started before the FE analysis Both programs should use the same WTBFunctions.dll at run time.

E.4.1. The Wind Coupling Process

Below is a brief description of the wind coupling algorithm implemented in Mechanical APDL:

- 1. Both Mechanical APDL and the aeroelastic code are run concurrently.
- 2. Both programs carry out data initialization separately.
- 3. At the beginning of a time step, set the aeroelastic code to active. The aeroelastic code computes and returns the turbine stiffness, damping and mass matrices to Mechanical APDL together with the aerodynamic force vector for the current time step. Put the aeroelastic code to sleep.
- 4. Mechanical APDL carries out the time integration to find the new solution at the end of the current time step.
- 5. The displacements, velocities and accelerations of the turbine freedoms (i.e. freedoms on the USER300 element) at the end of the step are stored to the shared common for the aeroelastic code.
- 6. Advance to next time step and return to step 3.

E.4.2. Data Exchange Routines

The following routines are used to facilitate data exchange between Mechanical APDL and the aeroelastic code as described above. They can be accessed from the aeroelastic code by linking in the

dynamic link library WTBFunctions.dll. Coding examples utilizing these routines in FORTRAN and C++ are available in the folder Program Files\Ansys Inc\v211\ansys\cus-tom\user\{platform}\Aeroelastic, where {platform} is a directory that uniquely identifies the hardware platform version: "Winx64" for 64-bit Windows.

```
SUBROUTINE GetWTBParamI(itype,id,ival,ierr)
!****
!****
         Routine gets an integer wind turbine parameter to common data area
!****
!****
         Arguments
!****
         itype (in ) Data type
!****
                       1 - Mechanical APDL run status
! * * * *
                            = -1 not started
!****
                            = 0 inactive (waiting)
!****
                           = 1 active (running)
! * * * *
                           = 2 stopped (finished/aborted)
!****
                       2 - Wind code run status
!****
                            = -1 not started
1 * * * *
                            = 0 inactive (waiting)
! * * * *
                            = 1 active (running)
!****
                              2 stopped (finished/aborted)
!****
                       {\it 3} - Number of active freedoms
!****
                       4 - Time step number
! * * * *
                 (in ) element identifier (currently unused, assume 1)
         id
!****
         ival
                 (out) parameter value
!****
         ierr
                 (out) exit code (0 if no error)
integer itype, id, ival, ierr
SUBROUTINE GetWTBParamR(itype,id,rval,ierr)
! * * * *
! * * * *
         Routine gets a real wind turbine parameter to common data area
! * * * *
!****
         Arguments
!****
         itype (in ) Data type
!****
                       1 - Analysis time
!****
                       2 - Time step
!****
         id
                 (in ) element identifier (currently unused, assume 1)
!****
         rval
                 (out) parameter value
         ierr
                 (out) exit code (0 if no error)
integer itype,id,ierr
double precision rval
SUBROUTINE GetWTBData(itype,id,array,narray,ierr)
! * * * *
!****
         Routine gets wind turbine data from common data area
! * * * *
! * * * *
         Arguments
!****
         itype (in ) Data type
1 * * * *
                       1 - Stif
!****
                       2 - Damp
! * * * *
                       3 - Mass
!****
                       4 - Load
!****
                       5 - Disp
! * * * *
                       6 - Velo
! * * * *
                       7 - Accn
!****
         id
                 (in ) element identifier (currently unused, assume 1)
!****
         array (out) data array
!****
         narray (i/o) size of array on input, actual array size on exit
!****
         ierr
                (out) exit code (0 if no error)
integer itype, id, narray, ierr
double precision array(*)
SUBROUTINE PutWTBParamI(itype,id,ival,ierr)
! * * * *
! * * * *
         Routine puts an integer wind turbine parameter to common data area
! * * * *
! * * * *
         Arguments
```

```
! * * * *
          itype (in ) Data type
! * * * *
                       1 - Mechanical APDL run status
!****
                           = -1 not started
!****
                            = 0 inactive (waiting)
!****
                           = 1 active (running)
! * * * *
                           = 2 stopped (finished/aborted)
! * * * *
                       2 - Wind code run status
!****
                           = -1 not started
! * * * *
                           = 0 inactive (waiting)
! * * * *
                           = 1 active (running)
!****
                           = 2 stopped (finished/aborted)
!****
                       3 - Number of active freedoms
! * * * *
                       4 - Time step number
!****
         id
                (in ) element identifier (currently unused, assume 1)
! * * * *
         ival (in ) parameter value
!****
         ierr (out) exit code (0 if no error)
integer itype, id, ival, ierr
SUBROUTINE PutWTBParamR(itype,id,rval,ierr)
!****
         Routine puts a real wind turbine parameter to common data area
! * * * *
!****
         Arguments
!****
         itype (in ) Data type
! * * * *
                       1 - Analysis time
                       2 - Time step
!****
!****
         id
                (in ) element identifier (currently unused, assume 1)
! * * * *
         rval (in ) parameter value
!****
         ierr (out) exit code (0 if no error)
integer itype, id, ierr
double precision rval
SUBROUTINE PutWTBData(itype,id,array,narray,ierr)
!****
         Routine puts wind turbine data to common data area
! * * * *
! * * * *
         Arguments
!****
         itype (in ) Data type
!****
                       1 - Stif
!****
                       2 - Damp
! * * * *
                       3 - Mass
!****
                       4 - Load
!****
                       5 - Disp
!****
                       6 - Velo
! * * * *
                       7 - Accn
!****
               (in ) element identifier (currently unused, assume 1)
         id
!****
         array (in ) data array
         narray (i/o) size of array on input, actual put size on exit
      ierr (out) exit code (0 if no error)
integer itype, id, narray, ierr
double precision array(*)
```

E.4.3. Important Analysis Notes

You must keep in mind the following when performing an aeroelastic analysis:

- After data initialization (e.g. data read in and checking, etc), the aeroelastic code should be put to sleep until the Mechanical APDL run status becomes inactive.
- The number of active freedoms is the number of freedoms in the aeroelastic model. This must be set up and put to the shared common by the aeroelastic code during the data initialization phase of the analysis.

- For the current usage, the aeroelastic code should always only put stiffness, damping, mass, and load data to the shared common, and get displacements, velocities, and accelerations from the shared common.
- The wind turbine array entries must correspond to the order of the element freedoms set up for the wind coupled USER300 element. Thus, freedoms 1 to 6 are UX, UY, UZ, ROTX, ROTY, and ROTZ freedoms of the interface node between the turbine and the support structure. The rest are generalized freedoms internal to the element. All the data must be stated in the structural coordinate axis system.
- The element matrices (i.e. stiffness, damping, and mass) are assumed to be given in packed symmetric form. The order of the packed symmetric matrix form in which the data are specified is defined as follows:

1	2	4	7	
	3	5	8	
		6	9	
	,		10	

- The units of the wind turbine data values are assumed to be consistent with the analysis units. No units conversion will be carried out by Mechanical APDL.
- It is assumed that identical time step sizes are used in both Mechanical APDL and the aeroelastic code. The solution times are controlled by data in Mechanical APDL since it is the one that solves the complete set of equations of the coupled system.
- The table below shows the explanation of the various exit code values (i.e. ierr):

Code	Meaning			
100	Invalid data type integer			
101	Specified array size too small to get			
102	Specified array size too big to put			
103	Number of active freedoms is unset			
104	Invalid array size specified			
201	Invalid run status specified			
202	Invalid number of active freedoms specified			
203	Invalid time step number specified			
301	Invalid time step value specified			

E.5. Example Analysis Using Provided "WindProg" Example for Aeroelastic Coupling

This example uses WindProg.exe, which represents a dummy aeroelastic analysis. Source code to allow you to compile this program is provided for both C++ and FORTRAN in {Installation Folder}\ansys\custom\user\{Platform}\Aeroelastic\WindDemos\{C++ | FORTRAN}. It assumes that the steps in Compiling a Custom Version of Mechanical APDL (p. 391) have been followed to create a custom executable for Mechanical APDL.

The example Mechanical APDL file is a simple line of pipe elements which is fully supported at node 6 and has the interface node positioned at node 1. The **WTBCREATE** macro is used to set up the user element and define the interface node.

To run the analysis, WindProg. exe must be able to find the WTBFunctions.dll either by having both files in the same folder, or by adding the folder containing WTBFunctions.dll to your Path system environment variable. First run WindProg and enter 3 as the number of freedoms, then run the following example within Mechanical APDL.

```
/FILNAME, wind02
/prep7
/TITLE, wind02, Wind coupling test
/com Wind coupling testing 02
antype, trans
nlgeom, off
et,1,pipe288
! define pipe section
secnum,1
sectype,1,pipe
secdata, 0.1, 0.02
MP, EX, 1,2.1e11
MP, PRXY, 1, 0.3
MP, ALPX, 1, 0.0
MP, DENS, 1, 7850.0
! define the tube
     1, 0.0, 0.0
n,
                1.0, 0.0
2.0, 0.0
        2,
n,
        3,
n,
                3.0, 0.0
       4,
n,
       5,
                4.0, 0.0
                5.0, 0.0
       6,
n,
type,1
mat,1
secnum,1
en, 1, 1, 2
en, 2, 2, 3
en, 3, 3, 4
en, 4, 4, 5
       5, 6
en, 5,
! define damping factors
alphad,0.3
betad, 0.001
! define turbine element
wtbcreate,,1,1 ! use Rayleigh damping
d. 6.all
finish
/SOLU
! CASE 1
F,1,FY,1.0e5
TINTP, 0.0
TIME, 1.0e-6
nsubst,1
solve
```

TIME, 0.1
nsubst, 10, 10, 10
OUTPR, all, 1
OUTRES, all, 1
solve
finish
/POST1
FORCE, STATIC
PRESOL, F
PRESOL, M
finish

E.6. References

The following references are cited in this appendix:

- 1. D. Kaufer et. al., Integrated Analysis of the Dynamics of Offshore Wind Trubines with Arbitrary Support Structures, Proc. of EWEC 2009, Marseille: EWEC, 2009.
- 2. M. Seidal et. Al., Validation of Offshore Load Simulations Using Measurement Data from the DOWNVInD Project, Proc. European Offshore Wind 2009, Stockholm, 2009.

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