

# Abaqus 6.10-EF

User Subroutines  
Reference Manual





# **Abaqus User Subroutines**

## **Reference Manual**

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## Preface

This section lists various resources that are available for help with using Abaqus Unified FEA software.

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# Contents

## 1. User Subroutines

### Abaqus/Standard subroutines

<b>CREEP:</b> Define time-dependent, viscoplastic behavior (creep and swelling).	1.1.1
<b>DFLOW:</b> Define nonuniform pore fluid velocity in a consolidation analysis.	1.1.2
<b>DFLUX:</b> Define nonuniform distributed flux in a heat transfer or mass diffusion analysis.	1.1.3
<b>DISP:</b> Specify prescribed boundary conditions.	1.1.4
<b>DLOAD:</b> Specify nonuniform distributed loads.	1.1.5
<b>FILM:</b> Define nonuniform film coefficient and associated sink temperatures for heat transfer analysis.	1.1.6
<b>FLOW:</b> Define nonuniform seepage coefficient and associated sink pore pressure for consolidation analysis.	1.1.7
<b>FRIC:</b> Define frictional behavior for contact surfaces.	1.1.8
<b>FRIC_COEF:</b> Define the frictional coefficient for contact surfaces.	1.1.9
<b>GAPCON:</b> Define conductance between contact surfaces or nodes in a fully coupled temperature-displacement analysis or pure heat transfer analysis.	1.1.10
<b>GAPELECTR:</b> Define electrical conductance between surfaces in a coupled thermal-electrical analysis.	1.1.11
<b>HARDINI:</b> Define initial equivalent plastic strain and initial backstress tensor.	1.1.12
<b>HETVAL:</b> Provide internal heat generation in heat transfer analysis.	1.1.13
<b>MPC:</b> Define multi-point constraints.	1.1.14
<b>ORIENT:</b> Provide an orientation for defining local material directions or local directions for kinematic coupling constraints or local rigid body directions for inertia relief.	1.1.15
<b>RSURFU:</b> Define a rigid surface.	1.1.16
<b>SDVINI:</b> Define initial solution-dependent state variable fields.	1.1.17
<b>SIGINI:</b> Define an initial stress field.	1.1.18
<b>UAMP:</b> Specify amplitudes.	1.1.19
<b>UANISOHYP<small>R</small> _INV:</b> Define anisotropic hyperelastic material behavior using the invariant formulation.	1.1.20
<b>UANISOHYP<small>R</small> _STRAIN:</b> Define anisotropic hyperelastic material behavior based on Green strain.	1.1.21
<b>UCORR:</b> Define cross-correlation properties for random response loading.	1.1.22
<b>UDMGINI:</b> Define the damage initiation criterion.	1.1.23
<b>UEL:</b> Define an element.	1.1.24
<b>UELMAT:</b> Define an element with access to materials.	1.1.25
<b>UEXPAN:</b> Define incremental thermal strains.	1.1.26
<b>UEXTERNALDB:</b> Manage user-defined external databases and calculate model-independent history information.	1.1.27

## CONTENTS

<b>UFIELD:</b> Specify predefined field variables.	1.1.28
<b>UFLUID:</b> Define fluid density and fluid compliance for hydrostatic fluid elements.	1.1.29
<b>UFLUIDLEAKOFF:</b> Define the fluid leak-off coefficients for pore pressure cohesive elements.	1.1.30
<b>UGENS:</b> Define the mechanical behavior of a shell section.	1.1.31
<b>UHARD:</b> Define the yield surface size and hardening parameters for isotropic plasticity or combined hardening models.	1.1.32
<b>UHYPERL:</b> Define a hypoelastic stress-strain relation.	1.1.33
<b>UHYPER:</b> Define a hyperelastic material.	1.1.34
<b>UINTER:</b> Define surface interaction behavior for contact surfaces.	1.1.35
<b>UMASFL:</b> Specify prescribed mass flow rate conditions for a convection/diffusion heat transfer analysis.	1.1.36
<b>UMAT:</b> Define a material's mechanical behavior.	1.1.37
<b>UMATHT:</b> Define a material's thermal behavior.	1.1.38
<b>UMESHMOTION:</b> Specify mesh motion constraints during adaptive meshing.	1.1.39
<b>UMOTION:</b> Specify motions during cavity radiation heat transfer analysis or steady-state transport analysis.	1.1.40
<b>UMULLINS:</b> Define damage variable for the Mullins effect material model.	1.1.41
<b>UPOREP:</b> Define initial fluid pore pressure.	1.1.42
<b>UPRESS:</b> Specify prescribed equivalent pressure stress conditions.	1.1.43
<b>UPSD:</b> Define the frequency dependence for random response loading.	1.1.44
<b>URDFIL:</b> Read the results file.	1.1.45
<b>USDFLD:</b> Redefine field variables at a material point.	1.1.46
<b>UTEMP:</b> Specify prescribed temperatures.	1.1.47
<b>UTRACLOAD:</b> Specify nonuniform traction loads.	1.1.48
<b>UTRS:</b> Define a reduced time shift function for a viscoelastic material.	1.1.49
<b>UVARM:</b> Generate element output.	1.1.50
<b>UWAVE:</b> Define wave kinematics for an analysis.	1.1.51
<b>VOIDRI:</b> Define initial void ratios.	1.1.52

### Abaqus/Explicit subroutines

<b>VDISP:</b> Specify prescribed boundary conditions.	1.2.1
<b>VDLOAD:</b> Specify nonuniform distributed loads.	1.2.2
<b>VFABRIC:</b> Define fabric material behavior.	1.2.3
<b>VFRIC:</b> Define frictional behavior for contact surfaces.	1.2.4
<b>VFRIC_COEF:</b> Define the frictional coefficient for contact surfaces.	1.2.5
<b>VFRICTION:</b> Define frictional behavior for contact surfaces.	1.2.6
<b>VUAMP:</b> Specify amplitudes.	1.2.7
<b>VUANISOHYPER_INV:</b> Define anisotropic hyperelastic material behavior using the invariant formulation.	1.2.8
<b>VUANISOHYPER_STRAIN:</b> Define anisotropic hyperelastic material behavior based on Green strain.	1.2.9
<b>VUEL:</b> Define an element.	1.2.10

<b>VUFIELD:</b> Specify predefined field variables.	1.2.11
<b>VUFLUIDEXCH:</b> Define the mass flow rate/heat energy flow rate for fluid exchange.	1.2.12
<b>VUFLUIDEXCHEFFAREA:</b> Define the effective area for fluid exchange.	1.2.13
<b>UVHARD:</b> Define the yield surface size and hardening parameters for isotropic plasticity or combined hardening models.	1.2.14
<b>VUINTER:</b> Define the interaction between contact surfaces.	1.2.15
<b>VUINTERACTION:</b> Define the contact interaction between surfaces with the general contact algorithm.	1.2.16
<b>VUMAT:</b> Define material behavior.	1.2.17
<b>VUSDFLD:</b> Redefine field variables at a material point.	1.2.18
<b>VUTRS:</b> Define a reduced time shift function for a viscoelastic material.	1.2.19
<b>VUVISCOSITY:</b> Define the shear viscosity for equation of state models.	1.2.20

## 2. Utility Routines

### Utility routines

Obtaining Abaqus environment variables	2.1.1
Obtaining the Abaqus job name	2.1.2
Obtaining the Abaqus output directory name	2.1.3
Obtaining parallel processes information	2.1.4
Obtaining part information	2.1.5
Obtaining material point information in an Abaqus/Standard analysis	2.1.6
Obtaining material point information in an Abaqus/Explicit analysis	2.1.7
Obtaining material point information averaged at a node	2.1.8
Obtaining node point information	2.1.9
Obtaining node to element connectivity	2.1.10
Obtaining stress invariants, principal stress/strain values and directions, and rotating tensors in an Abaqus/Standard analysis	2.1.11
Obtaining principal stress/strain values and directions in an Abaqus/Explicit analysis	2.1.12
Obtaining wave kinematic data in an Abaqus/Aqua analysis	2.1.13
Printing messages to the message or status file	2.1.14
Terminating an analysis	2.1.15
Obtaining sensor information	2.1.16
Accessing Abaqus materials	2.1.17
Accessing Abaqus thermal materials	2.1.18

## A. Index

User subroutines index	A.1
User subroutine functions listing	A.2



This manual describes all of the user subroutines and utility routines available in Abaqus. The interface and requirements for each user subroutine are discussed in detail. References to practical examples of most subroutines are also provided. Utility routines can be used within user subroutines to perform a variety of common tasks. The interface for all available utility routines appears in a separate chapter. For information on incorporating a user subroutine into an Abaqus analysis, see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

This manual is divided into three main sections:

- “Abaqus/Standard subroutines,” Section 1.1 covers all of the user subroutines available for use in an Abaqus/Standard analysis. Each section discusses a particular subroutine. The sections are organized alphabetically according to the subroutine name.
- “Abaqus/Explicit subroutines,” Section 1.2 covers all of the user subroutines available for use in an Abaqus/Explicit analysis. Each section discusses a particular subroutine. The sections are organized alphabetically according to the subroutine name.
- “Utility routines,” Section 2.1 covers all of the utility routines available for use in coding user subroutines. Each section discusses a task that can be performed using a utility routine. All of the utility routines associated with a particular task appear in the same section.



## 1. User Subroutines

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- “Abaqus/Standard subroutines,” Section 1.1
- “Abaqus/Explicit subroutines,” Section 1.2



## 1.1 Abaqus/Standard subroutines

- “CREEP,” Section 1.1.1
- “DFLOW,” Section 1.1.2
- “DFLUX,” Section 1.1.3
- “DISP,” Section 1.1.4
- “DLOAD,” Section 1.1.5
- “FILM,” Section 1.1.6
- “FLOW,” Section 1.1.7
- “FRIC,” Section 1.1.8
- “FRIC\_COEF,” Section 1.1.9
- “GAPCON,” Section 1.1.10
- “GAPELECTR,” Section 1.1.11
- “HARDINI,” Section 1.1.12
- “HETVAL,” Section 1.1.13
- “MPC,” Section 1.1.14
- “ORIENT,” Section 1.1.15
- “RSURFU,” Section 1.1.16
- “SDVINI,” Section 1.1.17
- “SIGINI,” Section 1.1.18
- “UAMP,” Section 1.1.19
- “UANISOHYPER\_INV,” Section 1.1.20
- “UANISOHYPER\_STRAIN,” Section 1.1.21
- “UCORR,” Section 1.1.22
- “UDMGINI,” Section 1.1.23
- “UEL,” Section 1.1.24
- “UELMAT,” Section 1.1.25
- “UEXPAN,” Section 1.1.26
- “UEXTERNALDB,” Section 1.1.27
- “UFIELD,” Section 1.1.28
- “UFLUID,” Section 1.1.29
- “UFLUIDLEAKOFF,” Section 1.1.30
- “UGENS,” Section 1.1.31
- “UHARD,” Section 1.1.32

- “UHYPEL,” Section 1.1.33
- “UHYPER,” Section 1.1.34
- “UINTER,” Section 1.1.35
- “UMASFL,” Section 1.1.36
- “UMAT,” Section 1.1.37
- “UMATHT,” Section 1.1.38
- “UMESHMOTION,” Section 1.1.39
- “UMOTION,” Section 1.1.40
- “UMULLINS,” Section 1.1.41
- “UPOREP,” Section 1.1.42
- “UPRESS,” Section 1.1.43
- “UPSD,” Section 1.1.44
- “URDFIL,” Section 1.1.45
- “USDFLD,” Section 1.1.46
- “UTEMP,” Section 1.1.47
- “UTRACLOAD,” Section 1.1.48
- “UTRS,” Section 1.1.49
- “UVARM,” Section 1.1.50
- “UWAVE,” Section 1.1.51
- “VOIDRI,” Section 1.1.52

## 1.1.1 CREEP: User subroutine to define time-dependent, viscoplastic behavior (creep and swelling).

**Product:** Abaqus/Standard

### References

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- “Rate-dependent plasticity: creep and swelling,” Section 20.2.4 of the Abaqus Analysis User’s Manual
- “Extended Drucker-Prager models,” Section 20.3.1 of the Abaqus Analysis User’s Manual
- “Modified Drucker-Prager/Cap model,” Section 20.3.2 of the Abaqus Analysis User’s Manual
- “Defining the gasket behavior directly using a gasket behavior model,” Section 29.6.6 of the Abaqus Analysis User’s Manual
- \*CAP CREEP
- \*CREEP
- \*DRUCKER PRAGER CREEP
- \*SWELLING
- “Verification of creep integration,” Section 3.2.6 of the Abaqus Benchmarks Manual

### Overview

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User subroutine **CREEP** will be called at all integration points of elements for which the material definition contains user-subroutine-defined metal creep, time-dependent volumetric swelling, Drucker-Prager creep, or cap creep behavior, during procedures that allow viscoplastic response of the above type to occur (such as the quasi-static procedure). This subroutine will also be called at all integration points of gasket elements for which the behavior definition contains user-subroutine-defined creep.

If user subroutine **CREEP** is used to define a material behavior, the subroutine:

- is intended to provide the “uniaxial” creep laws that are to be included in a general time-dependent, viscoplastic material formulation;
- can be used in the coupled-temperature displacement (“Fully coupled thermal-stress analysis,” Section 6.5.4 of the Abaqus Analysis User’s Manual), soils (“Coupled pore fluid diffusion and stress analysis,” Section 6.8.1 of the Abaqus Analysis User’s Manual), and quasi-static (“Quasi-static analysis,” Section 6.2.5 of the Abaqus Analysis User’s Manual) procedures;
- allows for the definition of creep laws for which the meaning and internal use depend on the material model with which they are being used;
- allows creep and swelling to be combined with rate-independent plastic behavior in a coupled manner, or they may simply be the only inelastic behaviors of the material, in which case Mises behavior is assumed;

## CREEP

- can use and update solution-dependent state variables; and
- can be used in conjunction with user subroutine **USDFLD** to redefine any field variables before they are passed in.

If user subroutine **CREEP** is used to define rate-dependent behavior in the thickness direction for a gasket, the subroutine:

- is intended to provide the creep laws that are used to prescribe the thickness-direction behavior for a gasket;
- can be used only in a quasi-static (“Quasi-static analysis,” Section 6.2.5 of the Abaqus Analysis User’s Manual) procedure;
- is used in a coupled form with the elastic-plastic model used to define the rate-independent part of the thickness-direction behavior of the gasket; and
- can use and update solution-dependent variables.

## Metals

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For metals whose material behavior includes metal creep and/or time-dependent volumetric swelling, the routine allows any “creep” and “swelling” laws (viscoplastic behavior) of the following general form to be defined:

$$\dot{\bar{\varepsilon}}^{cr} = g^{cr}(p, \tilde{q}, \bar{\varepsilon}^{sw}, \bar{\varepsilon}^{cr}, \text{time}, \dots),$$

$$\dot{\bar{\varepsilon}}^{sw} = g^{sw}(p, \tilde{q}, \bar{\varepsilon}^{sw}, \bar{\varepsilon}^{cr}, \text{time}, \dots),$$

where

- $\bar{\varepsilon}^{cr}$  is the uniaxial equivalent “creep” strain, conjugate to  $\tilde{q}$ , the Mises or Hill equivalent stress;  
 $\bar{\varepsilon}^{sw}$  is the volumetric swelling strain;  
 $p$  is the equivalent pressure stress,  $p = -\frac{1}{3}(\sigma_{11} + \sigma_{22} + \sigma_{33})$ ; and  
 $\tilde{q}$  is the equivalent deviatoric stress (Mises’ or, if anisotropic creep behavior is defined, Hill’s definition).

The user subroutine must define the increments of inelastic strain,  $\Delta\bar{\varepsilon}^{cr}$  and  $\Delta\bar{\varepsilon}^{sw}$ , as functions of  $p$  and  $\tilde{q}$  and any other variables used in the definitions of  $g^{cr}$  and  $g^{sw}$  (such as solution-dependent state variables introduced by you) and of the time increment,  $\Delta t$ . If any solution-dependent state variables are included in the definitions of  $g^{cr}$  and  $g^{sw}$ , they must also be integrated forward in time in this routine.

Abaqus computes the incremental creep strain (or the incremental viscoplastic strain) components as

$$\Delta\bar{\varepsilon}^{cr} = \frac{1}{3}\Delta\bar{\varepsilon}^{sw}\mathbf{R} + \Delta\bar{\varepsilon}^{cr}\mathbf{n},$$

where  $\mathbf{n}$  is the gradient of the deviatoric stress potential, defined as

$$\mathbf{n} = \frac{\partial \tilde{q}}{\partial \boldsymbol{\sigma}},$$

and  $\mathbf{R}$  is a matrix with the anisotropic swelling ratios in the diagonal if anisotropic swelling is defined; otherwise,  $\mathbf{R} = \mathbf{I}$ .

### Drucker-Prager materials

---

For materials that yield according to the extended Drucker-Prager plasticity models using Drucker-Prager creep, the routine allows any “creep” laws (viscoplastic behavior) of the following general form to be defined:

$$\dot{\bar{\varepsilon}}^{cr} = g^{cr}(\bar{\sigma}^{cr}, \bar{\varepsilon}^{cr}, \text{time}, \dots),$$

where

$\bar{\sigma}^{cr}$  is the equivalent creep stress defined as

$$\frac{q - p \tan \beta}{1 - \frac{1}{3} \tan \beta} \quad \text{if creep is defined in terms of uniaxial compression,}$$

$$\frac{q - p \tan \beta}{1 + \frac{1}{3} \tan \beta} \quad \text{if creep is defined in terms of uniaxial tension, and}$$

$$q - p \tan \beta \quad \text{if creep is defined in terms of pure shear,}$$

where  $q$  is the equivalent deviatoric Mises’ stress,  $p$  is the pressure stress, and  $\beta$  is the friction angle, and

$\bar{\varepsilon}^{cr}$  is the uniaxial equivalent “creep” strain, conjugate to  $\bar{\sigma}^{cr}$  such that  $\bar{\sigma}^{cr} \Delta \bar{\varepsilon}^{cr} = \sigma_{ij} \Delta \varepsilon_{ij}^{cr}$ .

The user subroutine must define the increment of inelastic strain,  $\Delta \bar{\varepsilon}^{cr}$ , as a function of  $\bar{\sigma}^{cr}$  and any other variables used in the definitions of  $g^{cr}$  (such as solution-dependent state variables introduced by you) and of the time increment,  $\Delta t$ . If any solution-dependent state variables are included in the definitions of  $g^{cr}$ , they must also be integrated forward in time in this routine.

Abaqus computes the incremental creep strain (or the incremental viscoplastic strain) components as

$$\Delta \varepsilon^{cr} = \frac{\Delta \bar{\varepsilon}^{cr}}{f^{cr}} \left( \frac{q}{\sqrt{(\epsilon \bar{\sigma})_0 \tan \psi)^2 + q^2}} \mathbf{n} + \frac{1}{3} \tan \psi \mathbf{I} \right),$$

where  $\mathbf{n} = \partial \tilde{q} / \partial \boldsymbol{\sigma}$ . The variable  $f^{cr}$  is determined in such a way that

$$f^{cr} = \frac{1}{\bar{\sigma}^{cr}} \boldsymbol{\sigma} : \frac{\partial G^{cr}}{\partial \boldsymbol{\sigma}},$$

and

$$G^{cr} = \sqrt{(\epsilon \bar{\sigma})_0 \tan \psi)^2 + q^2} - p \tan \psi$$

is the hyperbolic creep potential, where  $\psi(\theta, f^\alpha)$  is the dilation angle measured in the  $p$ - $q$  plane at high confining pressure,  $\bar{\sigma}|_0 = \bar{\sigma}|_{\dot{\varepsilon}^{pl}=0, \dot{\varepsilon}^{el}=0}$  is the initial yield stress, and  $\epsilon$  is the eccentricity. See “Extended Drucker-Prager models,” Section 20.3.1 of the Abaqus Analysis User’s Manual, for a discussion of  $\psi$ ,  $\epsilon$ , and  $\bar{\sigma}|_0$ .

### Capped Drucker-Prager materials

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For materials that yield according to the modified Drucker-Prager/Cap plasticity model using cap creep, the routine allows any “cohesion creep” and “consolidation creep” laws (viscoplastic behavior) of the following general form to be defined:

$$\dot{\varepsilon}_s^{cr} = g_s^{cr}(\bar{\sigma}^{cr}, \bar{\varepsilon}_s^{cr}, \text{time}, \dots),$$

$$\dot{\varepsilon}_c^{cr} = g_c^{cr}(\bar{p}^{cr}, \bar{\varepsilon}_c^{cr}, \text{time}, \dots),$$

where

$\bar{\sigma}^{cr}$  is the equivalent creep stress defined from uniaxial compression test data as

$$\bar{\sigma}^{cr} = \frac{q - p \tan \beta}{(1 - \frac{1}{3} \tan \beta)},$$

where  $q$  is the equivalent deviatoric Mises’ stress,  $p$  is the pressure stress, and  $\beta$  is the friction angle;

$\bar{\varepsilon}_s^{cr}$  is the equivalent cohesion creep uniaxial strain, conjugate to  $\bar{\sigma}^{cr}$  such that  $\bar{\sigma}^{cr} \Delta \bar{\varepsilon}_s^{cr} = \sigma : \Delta \varepsilon_s^{cr}$ , where  $\Delta \varepsilon_s^{cr}$  is defined below;

$\bar{p}^{cr} = p - p_a$  is the effective creep pressure ( $p = -\frac{1}{3}(\sigma_{11} + \sigma_{22} + \sigma_{33})$  and  $p_a$  is the cap hardening parameter); and

$\bar{\varepsilon}_c^{cr}$  is the volumetric consolidation creep strain.

The user subroutine must define the increments of inelastic strain,  $\Delta \bar{\varepsilon}_s^{cr}$  and/or  $\Delta \bar{\varepsilon}_c^{cr}$ , as functions of  $\bar{\sigma}^{cr}$  and/or  $\bar{p}^{cr}$  and any other variables used in the definitions of  $g_s^{cr}$  and  $g_c^{cr}$  (such as solution-dependent state variables introduced by you) and of the time increment,  $\Delta t$ . If any solution-dependent state variables are included in the definitions of  $g_s^{cr}$  and  $g_c^{cr}$ , they must also be integrated forward in time in this routine.

### Calculation of incremental creep strains for the cohesion mechanism

Abaqus computes the incremental creep strain (or the incremental viscoplastic strain) components of the cohesion mechanism as

$$\Delta \varepsilon_s^{cr} = \frac{\Delta \bar{\varepsilon}_s^{cr}}{f^{cr}} \left( \frac{q}{\sqrt{(0.1 \frac{d}{(1 - \frac{1}{3} \tan \beta)} \tan \beta)^2 + q^2}} \mathbf{n} + \frac{1}{3} \tan \beta \mathbf{I} \right),$$

where  $\mathbf{n} = \partial\tilde{q}/\partial\sigma$ ,  $d$  is the material cohesion, the variable  $f^{cr}$  is determined in such a way that

$$f^{cr} = \frac{1}{\bar{\sigma}^{cr}} \boldsymbol{\sigma} : \frac{\partial G_s^{cr}}{\partial \boldsymbol{\sigma}},$$

and  $G_s^{cr}$  is the cohesion creep potential

$$G_s^{cr} = \sqrt{(0.1 \frac{d}{(1 - \frac{1}{3} \tan \beta)} \tan \beta)^2 + q^2 - p \tan \beta}.$$

### Calculation of incremental creep strains for the consolidation mechanism

Abaqus computes the incremental creep strain (or the incremental viscoplastic strain) components of the consolidation mechanism as

$$\Delta \boldsymbol{\varepsilon}_c^{cr} = \frac{\Delta \bar{\boldsymbol{\varepsilon}}_c^{cr}}{G_c^{cr}} (R^2 q \mathbf{n} - \frac{1}{3}(p - p_a) \mathbf{I}),$$

where  $R$  controls the shape of the cap, and  $G_c^{cr}$  is the consolidation creep potential

$$G_c^{cr} = \sqrt{(p - p_a)^2 + (Rq)^2}.$$

Cohesion material properties are determined with a uniaxial compression test in which  $d\bar{\boldsymbol{\varepsilon}}_s^{cr} = \|d\boldsymbol{\varepsilon}_{11}^{cr}\|$ , and consolidation material properties are determined with a volumetric compression test in which  $d\bar{\boldsymbol{\varepsilon}}_c^{cr} = \|d\boldsymbol{\varepsilon}_{vol}^{cr}\|$ . Most likely,  $g_s^{cr}$  is a positive function of  $\bar{\sigma}^{cr}$ , and  $g_c^{cr}$  is a positive function of  $\bar{p}^{cr}$ .

---

### Gaskets

For gaskets whose behavior includes creep, the routine allows any “creep” law of the following general form to be defined:

$$\boldsymbol{\varepsilon}^{cr} = g^{cr}(\sigma, \boldsymbol{\varepsilon}^{cr}, \text{time}, \dots),$$

where  $\boldsymbol{\varepsilon}^{cr}$  is the compressive creep strain, conjugate to  $\sigma$ , the compressive stress in the gasket.

The user subroutine must define the increments of inelastic creep strain,  $\Delta \boldsymbol{\varepsilon}^{cr}$ , as functions of  $\sigma$  and any other variables used in the definitions of  $g^{cr}$  (such as solution-dependent state variables introduced by you) and of the time increment,  $\Delta t$ . If any solution-dependent state variables are included in the definitions of  $g^{cr}$ , they must also be integrated forward in time in this routine. Abaqus will automatically multiply this creep strain by the proper thickness (see “Defining the gasket behavior directly using a gasket behavior model,” Section 29.6.6 of the Abaqus Analysis User’s Manual) to obtain a creep closure.

---

### Integration schemes

Abaqus provides both explicit and implicit time integration of creep and swelling behavior defined in this routine. The choice of the time integration scheme depends on the procedure type, the procedure

definition, and whether a geometric linear or nonlinear analysis is requested (see “Rate-dependent plasticity: creep and swelling,” Section 20.2.4 of the Abaqus Analysis User’s Manual).

Implicit integration is generally more effective when the response period is long relative to typical relaxation times for the material. Simple high-temperature structural design applications usually do not need implicit integration, but more complicated problems (such as might arise in manufacturing processes), creep buckling applications, or nonstructural problems (such as geotechnical applications) often are integrated more efficiently by the implicit method provided in the program. If implicit integration is used with this subroutine, nonlinear equations must be solved at each time step and the variations of  $\Delta\bar{\varepsilon}^{cr}$ ,  $\Delta\bar{\varepsilon}^{sw}$ ,  $\Delta\bar{\varepsilon}_s^{cr}$ , or  $\Delta\bar{\varepsilon}_c^{cr}$  with respect to  $\bar{\varepsilon}^{cr}$ ,  $\bar{\varepsilon}^{sw}$ ,  $\bar{\varepsilon}_s^{cr}$ ,  $\bar{\varepsilon}_c^{cr}$ ,  $p$ ,  $\tilde{q}$ ,  $\bar{p}^{cr}$ , or  $\bar{\sigma}^{cr}$  must be defined in the subroutine. To obtain good convergence during implicit integration, it is essential to define these quantities accurately.

At the start of a new increment the subroutine is called once for each integration point to calculate the estimated creep strain based on the state at the start of the increment. Subsequently, it is called twice for each iteration if explicit integration is used: once to calculate the creep strain increment at the start of the increment and once to calculate it at the end of the increment. This is needed to test the validity of the time increment with respect to the user-specified maximum allowable difference in the creep strain increment. The flag **LEND** indicates whether the routine is called at the start or the end of the increment. The subroutine must use the corresponding values of time, temperature, field variables, and solution-dependent state variables in the calculation of the creep strain increment.

For implicit integration Abaqus uses a local iteration procedure to solve the nonlinear constitutive equations, and the subroutine is called multiple times. The exact number of calls depends on the convergence rate of the local iteration procedure and, hence, will vary from point to point. During these iterations it is possible for the values of the state variables to be far from their final values when the equations are solved. Therefore, the coding in the subroutine must adequately protect against arithmetic failures (such as floating point overflows) even when variables are passed in with physically unreasonable values. As in explicit integration, the variable **LEND** indicates whether the routine is called at the start or the end of the increment.

### **Constant stress assumption when defining creep and swelling**

---

When the creep and swelling behavior are defined by simple formulæ, it is often possible to calculate the increments of equivalent creep and swelling strain exactly if it is assumed that the stress is constant during the increment. This approach has the advantage that it provides very good accuracy within the constant stress assumption. It also avoids the problem that arises for some creep behavior definitions: that the creep strain rate becomes infinite at zero time (or strain). Otherwise, in such a case you must protect against causing arithmetic failures at the start of the solution.

### **Defining both plasticity and creep**

---

If both plasticity and creep are defined for a material, Abaqus will calculate the creep strain before entering the plasticity routines. The stresses passed into the creep routine may, therefore, exceed the yield stress.

## Interpretation of stress and strain variables

---

In finite-strain applications strain variables should be interpreted as logarithmic strains and stresses as “true” stress.

## User subroutine interface

---

```

SUBROUTINE CREEP (DECRA,DESWA,STATEV,SERD,EC,ESW,P,QTILD,
  1 TEMP,DTEMP,PREDEF,DPRED,TIME,DTIME,CMNAME,LEXIMP,LEND,
  2 COORDS,NSTATV,NOEL,NPT,LAYER,KSPT,KSTEP,KINC)
C
  INCLUDE 'ABA_PARAM.INC'
C
  CHARACTER*80 CMNAME
C
  DIMENSION DECRA(5),DESWA(5),STATEV(*),PREDEF(*),DPRED(*),
  1 TIME(2),EC(2),ESW(2),COORDS(*)

  user coding to define DECRA, DESWA

  RETURN
  END

```

## Variables to be defined

---

### In all cases

#### DECRA (1)

The definition depends on the usage:

- Metal creep:  $\Delta\bar{\varepsilon}^{cr}$ , equivalent (uniaxial) deviatoric creep strain increment.
- Drucker-Prager creep:  $\Delta\bar{\varepsilon}^{cr}$ , equivalent (uniaxial) creep strain increment.
- Capped Drucker-Prager creep:  $\Delta\bar{\varepsilon}_s^{cr}$ , equivalent (uniaxial) cohesion creep strain increment.
- Gasket creep:  $\Delta\varepsilon^{cr}$ , uniaxial compressive creep strain increment.

#### DESWA (1)

The definition depends on the usage:

- Metal creep:  $\Delta\bar{\varepsilon}^{sw}$ , volumetric swelling strain increment.
- Capped Drucker-Prager creep:  $\Delta\bar{\varepsilon}_c^{cr}$ , equivalent (volumetric) consolidation creep strain increment.
- Drucker-Prager and gasket creep: = 0.

**For implicit creep integration (`LEXIMP=1`, see below)****DECRA (2)**

The definition depends on the usage:

- Metal creep and Drucker-Prager creep:  $\partial\Delta\bar{\varepsilon}^{cr}/\partial\bar{\varepsilon}^{cr}$ .
- Capped Drucker-Prager creep:  $\partial\Delta\bar{\varepsilon}_s^{cr}/\partial\bar{\varepsilon}_s^{cr}$ .
- Gasket creep:  $\partial\Delta\varepsilon^{cr}/\partial\varepsilon^{cr}$ .

**DECRA (3)**

The definition depends on the usage:

- Metal creep:  $\partial\Delta\bar{\varepsilon}^{cr}/\partial\bar{\varepsilon}^{sw}$ .
- Drucker-Prager creep, gasket creep, and capped Drucker-Prager creep: = 0.

**DECRA (4)**

The definition depends on the usage:

- Metal creep:  $\partial\Delta\bar{\varepsilon}^{cr}/\partial p$ .
- Drucker-Prager creep, gasket creep, and capped Drucker-Prager creep: = 0.

**DECRA (5)**

The definition depends on the usage:

- Metal creep:  $\partial\Delta\bar{\varepsilon}^{cr}/\partial\tilde{q}$ .
- Drucker-Prager creep:  $\partial\Delta\bar{\varepsilon}^{cr}/\partial\bar{\sigma}^{cr}$ .
- Capped Drucker-Prager creep:  $\partial\Delta\bar{\varepsilon}_s^{cr}/\partial\bar{\sigma}^{cr}$ .
- Gasket creep:  $\partial\Delta\varepsilon^{cr}/\partial\sigma$ .

**DESWA (2)**

The definition depends on the usage:

- Metal creep:  $\partial\Delta\bar{\varepsilon}^{sw}/\partial\bar{\varepsilon}^{cr}$ .
- Drucker-Prager creep, gasket creep, and capped Drucker-Prager creep: = 0.

**DESWA (3)**

The definition depends on the usage:

- Metal creep:  $\partial\Delta\bar{\varepsilon}^{sw}/\partial\bar{\varepsilon}^{sw}$ .
- Capped Drucker-Prager creep:  $\partial\Delta\bar{\varepsilon}_c^{cr}/\partial\bar{\varepsilon}_c^{cr}$ .
- Drucker-Prager and gasket creep: = 0.

**DESWA (4)**

The definition depends on the usage:

- Metal creep:  $\partial\Delta\bar{\varepsilon}^{sw}/\partial p$ .
- Capped Drucker-Prager creep:  $\partial\Delta\bar{\varepsilon}_c^{cr}/\partial\bar{p}^{cr}$ .
- Drucker-Prager and gasket creep: = 0.

**DESWA (5)**

The definition depends on the usage:

- Metal creep:  $\partial \Delta \bar{\varepsilon}^{sw} / \partial \tilde{q}$ .
- Drucker-Prager creep, gasket creep, and capped Drucker-Prager creep: = 0.

**Variables that can be updated**

---

**STATEV**

An array containing the user-defined solution-dependent state variables at this point. This array will be passed in containing the values of these variables at the start of the increment unless they are updated in user subroutine **USDFLD** or **UEXPAN**, in which case the updated values are passed in. If any of the solution-dependent variables are being used in conjunction with the creep behavior and the routine was called at the end of the increment (**LEND=1**, see the definition of **LEND** below), they must be updated in this subroutine to their values at the end of the increment. Furthermore, if the solution-dependent state variables are defined as a function of the creep (swelling) strain increment, they must be updated based on the creep (swelling) strain increment computed as **EC (2) -EC (1)** (likewise **ESW (2) -ESW (1)**), where **EC (1)**, **EC (2)**, **ESW (1)**, and **ESW (2)** are defined below. You define the size of this array by allocating space for it (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information).

**SERD**

Magnitude of the strain energy rate density,  $\dot{W}$  (required only in  $C_t$ -integral calculations). The strain energy rate density is defined as

$$\dot{W} = \int_0^{\dot{\bar{\varepsilon}}^{cr}} q d\dot{\bar{\varepsilon}}^{cr} + \int_0^{\dot{\bar{\varepsilon}}^{sw}} p d\dot{\bar{\varepsilon}}^{sw} = \int_0^{\dot{\bar{\varepsilon}}} \boldsymbol{\sigma} : (d\dot{\bar{\varepsilon}}^{cr} + d\dot{\bar{\varepsilon}}^{sw}).$$

Elastic rates are ignored in the calculation of  $\dot{W}$ . The contour integral will, therefore, be path independent only for steady-state creep conditions; that is, when the creep straining dominates throughout the specimen.

**Variables passed in for information**

---

**EC (1)**

The definition depends on the usage:

- Metal creep and Drucker-Prager creep:  $\bar{\varepsilon}^{cr}$  at the start of the increment.
- Capped Drucker-Prager creep:  $\bar{\varepsilon}_s^{cr}$  at the start of the increment.
- Gasket creep:  $\varepsilon^{cr}$  at the start of the increment.

**EC (2)**

The definition depends on the usage:

- Metal creep and Drucker-Prager creep:  $\bar{\varepsilon}^{cr}$  at the end of the increment.

## CREEP

- Capped Drucker-Prager creep:  $\bar{\varepsilon}_s^{cr}$  at the end of the increment.
- Gasket creep:  $\varepsilon^{cr}$  at the end of the increment.

### ESW(1)

The definition depends on the usage:

- Metal creep:  $\bar{\varepsilon}^{sw}$  at the start of the increment.
- Capped Drucker-Prager creep:  $\bar{\varepsilon}_c^{cr}$  at the start of the increment.
- Drucker-Prager and gasket creep: = 0.

### ESW(2)

The definition depends on the usage:

- Metal creep:  $\bar{\varepsilon}^{sw}$  at the end of the increment.
- Capped Drucker-Prager creep:  $\bar{\varepsilon}_c^{cr}$  at the end of the increment.
- Drucker-Prager and gasket creep: = 0.

## P

The definition depends on the usage:

- Metal creep and Drucker-Prager creep:  $p = -\frac{1}{3}(\sigma_{11} + \sigma_{22} + \sigma_{33})$ , equivalent pressure stress (in soils analysis this is the equivalent effective pressure stress).
- Capped Drucker-Prager creep:  $\bar{p}^{cr} = p - p_a$ , effective creep pressure (in soils analysis  $p$  is the effective pressure stress).
- Gasket creep: = 0.

If **LEND=0**, the value is  $p$  or  $\bar{p}^{cr}$  at the beginning of the increment. If **LEND=1**, the value is  $p$  or  $\bar{p}^{cr}$  at the end of the increment.

## QTILD

The definition depends on the usage:

- Metal creep:  $\tilde{q}$ , Mises or Hill equivalent stress (the Hill formula is used if anisotropic creep is defined; see “Anisotropic creep” in “Rate-dependent plasticity: creep and swelling,” Section 20.2.4 of the Abaqus Analysis User’s Manual).
- Gasket creep:  $\sigma$ , the uniaxial compressive stress.
- Drucker-Prager creep:  $\bar{\sigma}^{cr}$ , equivalent creep stress (in soils analysis this is based on effective stresses).
- Capped Drucker-Prager creep:  $\bar{\sigma}^{cr}$ , equivalent creep stress (in soils analysis this is based on effective stresses).

If **LEND=0**, the value is  $\tilde{q}$  or  $\bar{\sigma}^{cr}$  at the beginning of the increment. If **LEND=1**, the value is  $\tilde{q}$  or  $\bar{\sigma}^{cr}$  at the end of the increment.

## TEMP

Temperature at the end of the increment.

**DTEMP**

Increment of temperature during the time increment.

**PREDEF**

An array containing the values of all of the user-specified predefined variables at this point at the end of the increment (initial values at the beginning of the analysis and current values during the analysis).

**DPRED**

An array containing the increments of all of the predefined variables during the time increment.

**TIME (1)**

Value of step time at the end of the increment.

**TIME (2)**

Value of total time at the end of the increment.

**DTIME**

Time increment.

**CMNAME**

User-specified material name or gasket behavior name, left justified. Some internal creep models are given names starting with the “ABQ\_” character string. To avoid conflict, you should not use “ABQ\_” as the leading string for **CMNAME**.

**LEXIMP**

Explicit/implicit flag.

If **LEXIMP=0**, explicit creep integration is being used and only **DECRA (1)** and **DESWA (1)** need be defined; **DECRA (I)** and **DESWA (I)**, **I=2,5**, need not be defined.

If **LEXIMP=1**, implicit creep integration is being used. The derivatives, **DECRA (I)** and **DESWA (I)**, **I=2,5**, should be defined accurately to achieve rapid convergence of the solution.

**LEND**

Start/end of increment flag.

If **LEND=0**, the routine is being called at the start of the increment. In this case **DECRA (1)** and **DESWA (1)** must be defined as the equivalent creep and swelling rates calculated at the beginning of the increment, multiplied by the time increment.

If **LEND=1**, the routine is being called at the end of the increment. In this case **DECRA (1)** and **DESWA (1)** must be defined as the equivalent creep and swelling rates calculated at the end of the increment, multiplied by the time increment. If applicable, the solution-dependent state variables **STATEV** must be updated as well.

**COORDS (3)**

An array containing the current coordinates of this point.

**NSTATV**

Number of solution-dependent state variables associated with this material or gasket behavior type (specified when space is allocated for the array; see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**KSTEP**

Step number.

**KINC**

Increment number.

---

**Example: Hyperbolic sine creep law**


---

Suppose that we wish to model a metal using the creep behavior

$$\dot{\varepsilon}^{cr} = A \left( \sinh \frac{q}{\sigma_0} \right)^n, \quad \dot{\varepsilon}^{sw} = 0,$$

where  $A$ ,  $\sigma_0$ , and  $n$  are constants.

User subroutine **CREEP** can be coded as follows:

```
SUBROUTINE CREEP (DECRA, DESWA, STATEV, SERD, EC, ESW, P, QTILD,
1 TEMP, DTEMP, PREDEF, DPRED, TIME, DTIME, CMNAME, LEXIMP, LEND,
2 COORDS, NSTATV, NOEL, NPT, LAYER, KSPT, KSTEP, KINC)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
C
DIMENSION DECRA(5), DESWA(5), STATEV(*), PREDEF(*), DPRED(*),
1 TIME(2), COORDS(*), EC(2), ESW(2)
C
C DEFINE CONSTANTS
C
```

```

A=
SIG0=
AN=
C
T1=EXP (QTILD/SIG0)
T2=EXP (-QTILD/SIG0)
DECRA (1) = A*(.5*(T1-T2))**AN*DTIME
IF (LEXIMP.EQ.1) THEN
    DECRA (5) = AN*A*(.5*(T1-T2))** (AN-1.) *DTIME/
1           SIG0*.5*(T1+T2)
END IF
C
RETURN
END

```

The derivative

$$\frac{\partial \Delta \bar{\varepsilon}^{cr}}{\partial q} = \frac{nA\Delta t}{\sigma_0} (\sinh \frac{q}{\sigma_0})^{n-1} \cosh \frac{q}{\sigma_0}$$

has been defined on the assumption that the subroutine will be used with implicit integration.



## 1.1.2 DFLOW: User subroutine to define nonuniform pore fluid velocity in a consolidation analysis.

**Product:** Abaqus/Standard

### References

---

- “Pore fluid flow,” Section 30.4.6 of the Abaqus Analysis User’s Manual
- \*DFLOW
- \*DSFLOW

### Overview

---

User subroutine **DFLOW**:

- can be used to define the variation of the seepage magnitude as a function of position, time, pore pressure, etc. in a soils consolidation analysis;
- will be called at each flow integration point for each element-based or surface-based nonuniform flow definition in the analysis; and
- ignores any amplitude references that may appear with the associated nonuniform flow definition.

### User subroutine interface

---

```

SUBROUTINE DFLOW(FLOW,U,KSTEP,KINC,TIME,NOEL,NPT,COORDS,
1 JLTYP,SNAME)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION TIME(2),COORDS(3)
CHARACTER*80 SNAME

```

*user coding to define FLOW*

```

RETURN
END

```

---

**Variable to be defined****FLOW**

Effective velocity of pore fluid crossing the surface at this point from the inside of the region modeled to the outside of the region modeled. Units are  $\text{LT}^{-1}$ . Effective velocity is the volumetric flow rate per unit area (refer to “Permeability,” Section 23.7.2 of the Abaqus Analysis User’s Manual).

**FLOW** will be passed into the routine as the magnitude of the seepage specified as part of the element-based or surface-based flow definition. If the magnitude is not defined, **FLOW** will be passed in as zero.

The effective velocity is not available for output purposes.

---

**Variables passed in for information****U**

Estimated pore pressure at this time at this point.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Current value of step time (defined only in transient analysis).

**TIME (2)**

Current value of total time (defined only in transient analysis).

**NOEL**

Element number.

**NPT**

Integration point number on the element’s surface.

**COORDS**

An array containing the coordinates of this point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**JLTYP**

Identifies the element face for which this call to **DFLOW** is being made through the element-based flow definition. This information is useful when several different nonuniform distributed flows are being imposed on an element at the same time. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for identification of element faces. The key is as follows:

<b>JLTYP</b>	<b>Flow type</b>
<b>0</b>	Surface-based load
<b>11</b>	S1NU
<b>12</b>	S2NU
<b>13</b>	S3NU
<b>14</b>	S4NU
<b>15</b>	S5NU
<b>16</b>	S6NU

**SNAME**

Surface name for which this call to **DFLOW** is being made through the surface-based flow definition (**JLTYP**=0). For an element-based flow definition the surface name is passed in as a blank.



### 1.1.3 DFLUX: User subroutine to define nonuniform distributed flux in a heat transfer or mass diffusion analysis.

**Product:** Abaqus/Standard

#### References

---

- “Thermal loads,” Section 30.4.4 of the Abaqus Analysis User’s Manual
- “Mass diffusion analysis,” Section 6.9.1 of the Abaqus Analysis User’s Manual
- \*DFLUX
- \*DSFLUX
- “DFLUX,” Section 4.1.1 of the Abaqus Verification Manual

#### Overview

---

User subroutine **DFLUX**:

- can be used to define a nonuniform distributed flux as a function of position, time, temperature, element number, integration point number, etc. in a heat transfer or mass diffusion analysis;
- will be called at each flux integration point for each element-based or surface-based (heat transfer only) nonuniform distributed flux definition in the analysis;
- ignores any amplitude references that may appear with the associated nonuniform distributed flux definition; and
- uses the nodes as flux integration points for first-order heat transfer, first-order coupled temperature-displacement, and mass diffusion elements.

#### User subroutine interface

---

```
SUBROUTINE DFLUX(FLUX,SOL,KSTEP,KINC,TIME,NOEL,NPT,COORDS,
  1 JLTYP,TEMP,PRESS,SNAME)
C
  INCLUDE 'ABA_PARAM.INC'
C
  DIMENSION FLUX(2), TIME(2), COORDS(3)
  CHARACTER*80 SNAME
```

*user coding to define FLUX(1) and FLUX(2)*

```
RETURN
END
```

**Variables to be defined**

---

**FLUX (1)**

Magnitude of flux flowing into the model at this point. In heat transfer cases the units are  $JT^{-1}L^{-2}$  for surface fluxes and  $JT^{-1}L^{-3}$  for body flux. In transient heat transfer cases where a non-default amplitude is used to vary the applied fluxes, the time average flux over the time increment must be defined rather than the value at the end of the time increment. In mass diffusion cases the units are  $PLT^{-1}$  for surface fluxes and  $PT^{-1}$  for body flux.

**FLUX (1)** will be passed into the routine as the magnitude of the flux specified as part of the element-based or surface-based flux definition. If the magnitude is not defined, **FLUX (1)** will be passed in as zero.

This flux is not available for output purposes.

**FLUX (2)**

In heat transfer cases:  $dq/d\theta$ , the rate of change of the flux with respect to the temperature at this point. The units are  $JT^{-1}L^{-2}\theta^{-1}$  for surface fluxes and  $JT^{-1}L^{-3}\theta^{-1}$  for body flux.

In mass diffusion cases:  $dq/dc$ , the rate of change of the flux with respect to the mass concentration at this point. The units are  $LT^{-1}$  for surface fluxes and  $T^{-1}$  for body flux.

The convergence rate during the solution of the nonlinear equations in an increment is improved by defining this value, especially when the flux is a strong function of temperature in heat transfer analysis or concentration in mass diffusion analysis.

**Variables passed in for information**

---

**SOL**

Estimated value of the solution variable (temperature in a heat transfer analysis or concentration in a mass diffusion analysis) at this time at this point.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Current value of step time (defined only in transient analysis).

**TIME (2)**

Current value of total time (defined only in transient analysis).

**NOEL**

Element number.

**NPT**

Integration point number in the element or on the element's surface. The integration scheme depends on whether this is a surface or a body flux.

**COORDS**

An array containing the coordinates of this point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**JLTYP**

Identifies the flux type for which this call to **DFLUX** is being made. The flux type may be a body flux, a surface-based flux, or an element-based surface flux. For element-based surface fluxes, this variable identifies the element face for which this call to **DFLUX** is being made. This information is useful when several different nonuniform distributed fluxes are being imposed on an element at the same time. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for element face identification. The key is as follows:

<b>JLTYP</b>	<b>Flux type</b>
<b>0</b>	Surface-based flux
<b>1</b>	BFNU
<b>11</b>	S1NU (SNEGNU for heat transfer shells)
<b>12</b>	S2NU (SPOSNU for heat transfer shells)
<b>13</b>	S3NU
<b>14</b>	S4NU
<b>15</b>	S5NU
<b>16</b>	S6NU

**TEMP**

Current value of temperature at this integration point (defined only for a mass diffusion analysis). Temperature for a heat transfer analysis is passed in as variable **SOL**.

**PRESS**

Current value of the equivalent pressure stress at this integration point (defined only for a mass diffusion analysis).

**SNAME**

Surface name for a surface-based flux definition (**JLTYP**=0). For a body flux or an element-based surface flux the surface name is passed in as blank.



## 1.1.4 DISP: User subroutine to specify prescribed boundary conditions.

**Product:** Abaqus/Standard

### References

---

- “Boundary conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.3.1 of the Abaqus Analysis User’s Manual
- “Connector actuation,” Section 28.1.3 of the Abaqus Analysis User’s Manual
- \*BOUNDARY
- \*CONNECTOR MOTION
- “Riser dynamics,” Section 11.1.2 of the Abaqus Example Problems Manual
- “DISP,” Section 4.1.2 of the Abaqus Verification Manual
- “\*BOUNDARY,” Section 5.1.4 of the Abaqus Verification Manual

### Overview

---

User subroutine **DISP**:

- can be used to define the magnitudes of prescribed boundary conditions or connector motions;
- is called for all degrees of freedom listed in a user-subroutine-defined boundary condition or connector motion definition;
- redefines any magnitudes that may be specified (and possibly modified by an amplitude) as part of the associated boundary condition or connector motion definition; and
- ignores the specified type, if any, of the associated boundary condition or connector motion definition.

### User subroutine interface

---

```
SUBROUTINE DISP(U,KSTEP,KINC,TIME,NODE,NOEL,JDOF,COORDS)
C
      INCLUDE 'ABA_PARAM.INC'
C
      DIMENSION U(3),TIME(2),COORDS(3)
C
```

*user coding to define U*

```
RETURN
END
```

---

**Variable to be defined****U(1)**

Total value of the prescribed variable at this point. The variable may be displacement, rotation, pore pressure, temperature, etc., depending on the degree of freedom constrained. **U(1)** will be passed into the routine as the value defined by any magnitude and/or amplitude specification for the boundary condition or connector motion.

If the analysis procedure requires that the time derivatives of prescribed variables be defined (for example, in a dynamic analysis the velocity and acceleration, as well as the value of the variable, are needed),  $du/dt$  must be given in **U(2)** and  $d^2u/dt^2$  in **U(3)**. The total value of the variable and its time derivatives must be given in user subroutine **DISP**, regardless of the type of boundary condition or connector motion.

---

**Variables passed in for information****KSTEP**

Step number.

**KINC**

Increment number.

**TIME(1)**

Current value of step time.

**TIME(2)**

Current value of total time.

**NODE**

Node number. This variable cannot be used if user subroutine **DISP** is used to prescribe connector motions.

**NOEL**

Element number. This variable cannot be used if user subroutine **DISP** is used to prescribe boundary conditions.

**JDOF**

Degree of freedom.

**COORDS**

An array containing the current coordinates of this point. These are the coordinates at the end of the prior increment if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the node. This array cannot be used if user subroutine **DISP** is used to prescribe connector motions.

## 1.1.5 DLOAD: User subroutine to specify nonuniform distributed loads.

**Product:** Abaqus/Standard

### References

---

- “Distributed loads,” Section 30.4.3 of the Abaqus Analysis User’s Manual
- \*DLOAD
- \*DSLOAD
- “Nonuniform crack-face loading and  $J$ -integrals,” Section 1.16.7 of the Abaqus Benchmarks Manual
- “Pure bending of a cylinder: CAXA elements,” Section 1.3.33 of the Abaqus Verification Manual
- “Cylinder subjected to asymmetric pressure loads: CAXA elements,” Section 1.3.35 of the Abaqus Verification Manual
- “Patch test for axisymmetric elements,” Section 1.5.4 of the Abaqus Verification Manual
- “Transient internal pressure loading of a viscoelastic cylinder,” Section 2.2.8 of the Abaqus Verification Manual
- “DLOAD,” Section 4.1.3 of the Abaqus Verification Manual

### Overview

---

User subroutine **DLOAD**:

- can be used to define the variation of the distributed load magnitude as a function of position, time, element number, load integration point number, etc.;
- will be called at each load integration point for each element-based or surface-based nonuniform distributed load definition during stress analysis;
- will be called at each stiffness integration point for computing the effective axial force, ESF1, for pipe elements subjected to nonuniform load types PENU and PINU;
- cannot be used in mode-based procedures to describe the time variation of the load; and
- ignores any amplitude references that may appear with the associated step definition or nonuniform distributed load definition.

### User subroutine interface

---

```

SUBROUTINE DLOAD (F,KSTEP,KINC,TIME,NOEL,NPT,LAYER,KSPT,
1 COORDS,JLTYP,SNAME)
C
INCLUDE 'ABA_PARAM.INC'
C

```

## DLOAD

```
DIMENSION TIME(2), COORDS (3)
CHARACTER*80 SNAME
```

*user coding to define **F***

```
RETURN
END
```

### Variable to be defined

---

#### **F**

Magnitude of the distributed load. Units are  $FL^{-2}$  for surface loads and  $FL^{-3}$  for body forces. **F** will be passed into the routine as the magnitude of the load specified as part of the element-based or surface-based distributed load definition. If the magnitude is not defined, **F** will be passed in as zero. For a static analysis that uses the modified Riks method (“Static stress analysis,” Section 6.2.2 of the Abaqus Analysis User’s Manual) **F** must be defined as a function of the load proportionality factor,  $\lambda$ . The distributed load magnitude is not available for output purposes.

### Variables passed in for information

---

#### **KSTEP**

Step number.

#### **KINC**

Increment number.

#### **TIME (1)**

Current value of step time or current value of the load proportionality factor,  $\lambda$ , in a Riks step.

#### **TIME (2)**

Current value of total time.

#### **NOEL**

Element number.

#### **NPT**

Load integration point number within the element or on the element’s surface, depending on the load type. (Stiffness integration point number while computing effective axial force, ESF1, for pipe elements subjected to load types PENU and PINU.)

#### **LAYER**

Layer number (for body forces in layered solids).

#### **KSPT**

Section point number within the current layer.

**COORDS**

An array containing the coordinates of the load integration point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point. For axisymmetric elements that allow nonaxisymmetric deformation, **COORDS (3)** is the angular position of the integration point, in degrees.

**JLTYP**

Identifies the load type for which this call to **DLOAD** is being made. The load type may be a body force, a surface-based load, or an element-based surface load. For element-based surface loads, this variable identifies the element face for which this call to **DLOAD** is being made. This information is useful when several different nonuniform distributed loads are being imposed on an element at the same time. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for element face identification. The key is as follows:

<b>JLTYP</b>	<b>Load type</b>
0	Surface-based load
1	BXNU
1	BRNU
2	BYNU (except for axisymmetric elements)
2	BZNU (for axisymmetric elements only)
3	BZNU (for three-dimensional elements and asymmetric-axisymmetric elements)
20	PNU
21	P1NU
22	P2NU
23	P3NU
24	P4NU
25	P5NU
26	P6NU
27	PINU
28	PENU
41	PXNU
42	PYNU
43	PZNU

## DLOAD

### SNAME

Surface name for a surface-based load definition (**JLTYP**=0). For a body force or an element-based surface load the surface name is passed in as blank.

## 1.1.6 FILM: User subroutine to define nonuniform film coefficient and associated sink temperatures for heat transfer analysis.

**Product:** Abaqus/Standard

### References

---

- “Thermal loads,” Section 30.4.4 of the Abaqus Analysis User’s Manual
- \*CFILM
- \*FILM
- \*SFILM
- “Temperature-dependent film condition,” Section 1.3.41 of the Abaqus Verification Manual

### Overview

---

User subroutine **FILM**:

- can be used to define a node-based, element-based, or surface-based nonuniform film coefficient;
- can be used to define sink temperatures as functions of position, time, temperature, node number, element number, integration point number, etc.;
- will be called during procedures that allow heat transfer analysis at each node or surface integration point of those surfaces and elements for which node-based, element-based, or surface-based nonuniform film conditions are defined;
- ignores any amplitude references for the sink temperature or film coefficient that may appear with the associated nonuniform film definition; and
- uses the nodes for first-order heat transfer elements as surface integration points for both element-based and surface-based films.

### User subroutine interface

---

```

SUBROUTINE FILM(H,SINK,TEMP,KSTEP,KINC,TIME,NOEL,NPT,
1 COORDS,JLTYP,FIELD,NFIELD,SNAME,NODE,AREA)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION H(2),TIME(2),COORDS(3), FIELD(NFIELD)
CHARACTER*80 SNAME

```

*user coding to define H(1), H(2), and SINK*

```

RETURN
END

```

**Variables to be defined**

---

**H(1)**

Film coefficient at this point. Units are  $JT^{-1}L^{-2}\theta^{-1}$ . **H(1)** will be passed into the routine as the magnitude of the film coefficient specified as part of the node-based, element-based, or surface-based film condition definition. If the magnitude is not defined, **H(1)** will be initialized to zero. This film coefficient is not available for output purposes.

**H(2)**

$dh/d\theta$ , rate of change of the film coefficient with respect to the surface temperature at this point. Units are  $JT^{-1}L^{-2}\theta^{-2}$ . The rate of convergence during the solution of the nonlinear equations in an increment is improved by defining this value, especially when the film coefficient is a strong function of surface temperature.

**SINK**

Sink temperature. **SINK** will be passed into the routine as the sink temperature specified as part of the node-based, element-based, or surface-based film condition definition. If the sink temperature is not defined, **SINK** will be initialized to zero. This sink temperature is not available for output purposes.

**Variables passed in for information**

---

**TEMP**

Estimated surface temperature at this time at this point.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME(1)**

Current value of step time.

**TIME(2)**

Current value of total time.

**NOEL**

Element number. This variable is passed in as zero for node-based films.

**NPT**

Surface integration point number. This variable is passed in as zero for node-based films.

**COORDS**

An array containing the coordinates of this point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**JLTYP**

Identifies the element face for which this call to **FILM** is being made for an element-based film coefficient specification. This information is useful when several different nonuniform film conditions are being imposed on an element at the same time. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for element face identification. The key is as follows:

<b>JLTYP</b>	<b>Film type</b>
<b>0</b>	Node-based or surface-based loading
<b>11</b>	F1NU (FNEGNU for heat transfer shells)
<b>12</b>	F2NU (FPOSNU for heat transfer shells)
<b>13</b>	F3NU
<b>14</b>	F4NU
<b>15</b>	F5NU
<b>16</b>	F6NU

**FIELD**

Interpolated values of field variables at this point.

**NFIELD**

Number of field variables.

**SNAME**

Surface name for which this call to **FILM** is being made for a surface-based film coefficient specification (**JLTYP**=0). This variable is passed in as blank for both node-based and element-based films.

**NODE**

Node number. This variable is passed in as zero for both element-based and surface-based films.

**AREA**

Nodal area for node-based films. **AREA** will be passed into the routine as the nodal area specified as part of the node-based film coefficient specification. This nodal area is not available for output purposes. This variable is passed in as zero for both element-based and surface-based films.



- 1.1.7 FLOW: User subroutine to define nonuniform seepage coefficient and associated sink pore pressure for consolidation analysis.**

**Product:** Abaqus/Standard

## References

---

- “Pore fluid flow,” Section 30.4.6 of the Abaqus Analysis User’s Manual
- \*FLOW
- \*SFLOW

## Overview

---

User subroutine **FLOW**:

- can be used in a soils consolidation analysis to define the variation of the reference pore pressure and the seepage coefficient as functions of position, time, pore pressure, element number, integration point number, etc.;
- will be called at each integration point of element surfaces for which element-based or surface-based nonuniform surface seepage flow is defined; and
- ignores any amplitude references that may appear with the associated nonuniform flow definition.

## User subroutine interface

---

```

SUBROUTINE FLOW(H,SINK,U,KSTEP,KINC,TIME,NOEL,NPT,COORDS,
1 JLTYP,SNAME)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION TIME(2), COORDS(3)
CHARACTER*80 SNAME

```

*user coding to define H and SINK*

```

RETURN
END

```

---

**Variables to be defined****H**

Seepage coefficient at this point. Units are  $F^{-1}L^3T^{-1}$ . **H** will be passed into the routine as the reference seepage coefficient value specified as part of the element-based or surface-based flow definition. If the reference value is not defined, **H** will be passed in as zero.

**SINK**

Sink pore pressure. **SINK** will be passed into the routine as the reference pore pressure value specified as part of the element-based or surface-based flow definition. If the reference value is not defined, **SINK** will be passed in as zero.

---

**Variables passed in for information****U**

Estimated surface total pore pressure at this time and at this point.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Current value of step time (defined only in transient analysis).

**TIME (2)**

Current value of total time (defined only in transient analysis).

**NOEL**

Element number.

**NPT**

Surface integration point number.

**COORDS**

An array containing the coordinates of this integration point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**JLTYP**

Identifies the element face for which this call to **FLOW** is being made for an element-based flow. This information is useful when several nonuniform flow conditions are being imposed on an element at the same time. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for identification of the element faces. The key is as follows:

<b>JLTYP</b>	<b>Flow type</b>
<b>0</b>	Surface-based flow
<b>61</b>	Q1NU
<b>62</b>	Q2NU
<b>63</b>	Q3NU
<b>64</b>	Q4NU
<b>65</b>	Q5NU
<b>66</b>	Q6NU

**SNAME**

Surface name for which this call to **FLOW** is being made for a surface-based flow (**JLTYP**=0). For an element-based flow the surface name is passed in as a blank.



## 1.1.8 FRIC: User subroutine to define frictional behavior for contact surfaces.

**Product:** Abaqus/Standard

### References

---

- “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual
- \*FRICTION
- “Thermal-stress analysis of a disc brake,” Section 5.1.1 of the Abaqus Example Problems Manual
- “**FRIC**,” Section 4.1.4 of the Abaqus Verification Manual

### Overview

---

User subroutine **FRIC**:

- can be used to define the frictional behavior between contacting surfaces;
- can be used when the extended versions of the classical Coulomb friction model provided in Abaqus are too restrictive and a more complex definition of shear transmission between contacting surfaces is required;
- will be called at points on the slave surface of a contact pair and at the integration points in a contact element (only when the contact point is closed) for which the contact interaction property model contains user-subroutine-defined friction;
- must provide the entire definition of shear interaction between the contacting surfaces; and
- can use and update solution-dependent state variables.

### User subroutine interface

---

```

SUBROUTINE FRIC (LM, TAU, DDTDDG, DDTDDP, DSLIP, SED, SFD,
1 DDTDDT, PNEWDT, STATEV, DGAM, TAULM, PRESS, DPRESS, DDPDDH, SLIP,
2 KSTEP, KINC, TIME, DTIME, NOEL, CINAME, SLNAME, MSNAME, NPT, NODE,
3 NPATCH, COORDS, RCOORD, DROT, TEMP, PREDEF, NFDIR, MCRD, NPRED,
4 NSTATV, CHRLNGTH, PROPS, NPROPS)
C
      INCLUDE 'ABA_PARAM.INC'
C
      CHARACTER*80 CINAME, SLNAME, MSNAME
C
      DIMENSION TAU(NFDIR), DDTDDG(NFDIR, NFDIR), DDTDDP(NFDIR),
1      DSLIP(NFDIR), DDTDDT(NFDIR, 2), STATEV(*), DGAM(NFDIR),
2      TAULM(NFDIR), SLIP(NFDIR), TIME(2), COORDS(MCRD),
3      RCOORD(MCRD), DROT(2, 2), TEMP(2), PREDEF(2, *), PROPS(NPROPS)

```

*user coding to define **LM**, **TAU**, **DDTDDG**, **DDTDDP**,  
and, optionally, **DSLIP**, **SED**, **SFD**, **DDTDDT**, **PNEWDT**, **STATEV***

```
RETURN
END
```

## Variables to be defined

---

### In all cases

#### **LM**

Relative motion flag. User subroutine **FRIC** is called only if the contact point is determined to be closed; that is, if the contact pressure is positive (the contact point was closed in the previous iteration) or if the contact point is overclosed (the contact point was open in the previous iteration).

During iterations **LM** is passed into the subroutine as the value defined during the previous iteration. At the start of an increment or if the contact point opened during the previous iteration, this variable will be passed into the routine depending on the contact condition in the previous increment. If the contact point was slipping, **LM** is equal to 0; if the contact point was sticking, **LM** is equal to 1; and if the contact point was open, **LM** is equal to 2.

Set **LM** equal to 0 if relative motion is allowed (either due to slip or elastic stick). In this case the subroutine must specify the frictional stress  $\tau_1$  (and  $\tau_2$  for three-dimensional analysis) as a function of the relative sliding motion  $\gamma_1$  (and  $\gamma_2$ ), the interface contact pressure  $p$ , and other predefined or user-defined state variables. In addition, the subroutine must define the derivatives of the frictional stress with respect to  $\gamma_1$ , ( $\gamma_2$ ), and  $p$ . For instance, in the case of isotropic elastic sticking,  $\partial\tau_1/\partial\gamma_1 = \partial\tau_2/\partial\gamma_2 = k_{elas}$ ,  $\partial\tau_1/\partial\gamma_2 = \partial\tau_2/\partial\gamma_1 = 0$ , where  $k_{elas}$  is the elastic stiffness of the interface.

Set **LM** equal to 1 if no relative motion is allowed; a rigid sticking condition at the interface is enforced by a Lagrange multiplier method. In this case no further variables need to be updated. If **LM** is always set to 1, a “perfectly rough” interface is created. It is not advisable to set **LM** to 1 when the finite-sliding, surface-to-surface contact formulation is used.

Set **LM** equal to 2 if friction is ignored (frictionless sliding is assumed). In this case no further variables need to be updated. If **LM** is always set to 2, a “perfectly smooth” interface is created.

You can make decisions about the stick/slip condition based on incremental slip information and calculated frictional stresses. These quantities are passed in by Abaqus/Standard, as discussed below.

To avoid convergence problems for the general class of frictional contact problems, set **LM** to 2 and exit this routine if the contact point was open at the end of the previous increment; that is, if Abaqus/Standard sets **LM**=2 when it calls this routine, simply exit the routine.

## If the return value of **LM** is 0

### **TAU (NFDIR)**

These values are passed in as the values of the frictional stress components,  $\tau_\alpha$ , at the beginning of the increment and must be updated to the values at the end of the increment. Here, and in the rest of this description, Greek subscripts ( $\alpha, \beta$ ) refer to frictional shear directions. The orientation of these directions on contact surfaces is defined in “Contact formulations in Abaqus/Standard,” Section 34.1.1 of the Abaqus Analysis User’s Manual.

### **DDTDDG (NFDIR, NFDIR)**

$\partial\Delta\tau_\alpha/\partial\Delta\gamma_\beta$ , partial derivative of the frictional stress in direction  $\alpha$  with respect to the relative motion in direction  $\beta$ .

### **DDTDDP (NFDIR)**

$\partial\Delta\tau_\alpha/\partial\Delta p$ , partial derivative of the frictional stress in direction  $\alpha$  with respect to the contact pressure. Since these terms yield an unsymmetric contribution to the stiffness matrix, they are used only if the unsymmetric equation solver is used (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual).

---

## Variables that can be updated

### **DSLIP (NFDIR)**

$\Delta\gamma_\alpha^{sl}$ , increment in nonrecoverable sliding motion (slip). If **LM** was 0 in the previous iteration, this array is passed in as the user-defined values during the previous iteration; otherwise, it will be zero. The array should be updated only if the return value of **LM** is 0.

This array is useful to detect slip reversals between iterations. It is used by the output options to indicate whether this point is sticking or slipping. Upon convergence of an increment, the values in **DSLIP (NFDIR)** are accumulated in **SLIP (NFDIR)**, which are stored as the plastic strains.

### **SED**

This variable is passed in as the value of the elastic energy density at the start of the increment and should be updated to the elastic energy density at the end of the increment. This variable is used for output only and has no effect on other solution variables.

### **SFD**

This variable should be defined as the incremental frictional dissipation. The units are energy per unit area if the contact element or contact pair calling **FRIC** uses stresses as opposed to forces. For regular stress analysis this variable is used for output only and has no effect on other solution variables. In coupled temperature-displacement analysis the dissipation is converted into heat if the gap heat generation model is used. If **SFD** is not defined, the heat generation is calculated based on the dissipation obtained as the product of the slip increment, **DSLIP**, and the frictional stress, **TAU**.

**DDTDDT (NFDIR, 2)**

$\partial\Delta\tau_\alpha/\partial\Delta\theta_1$ ,  $\partial\Delta\tau_\alpha/\partial\Delta\theta_2$  partial derivatives of the frictional stress in direction  $\alpha$  with respect to the temperatures of the two surfaces. This is required only for coupled temperature-displacement elements, in which the frictional stress is a function of the surface temperatures.

**PNEWDT**

Ratio of suggested new time increment to the time increment currently being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **FRIC**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** greater than 1.0 will be ignored and values of **PNEWDT** less than 1.0 will cause the job to terminate.

**STATEV (NSTATV)**

An array containing the user-defined solution-dependent state variables. You specify the number of available state variables; see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for details. This array will be passed in containing the values of these variables at the start of the increment. If any of the solution-dependent state variables is being used in conjunction with the friction behavior, they must be updated in this subroutine to their values at the end of the increment.

---

**Variables passed in for information**
**DGAM (NFDIR)**

If **LM** was set to 0 in the previous iteration, this value is the increment of sliding motion in the current increment,  $\Delta\gamma_\alpha$ . Otherwise, it will be zero. Comparison with **DSLIP (NFDIR)** makes it possible to determine whether slip changes to stick at this point and/or if there is a slip direction reversal occurring at this point.

**TAULM (NFDIR)**

If **LM** was set to 1 in the previous iteration, this value is the current value of the constraint stress at the end of the increment,  $\tau_\alpha^{LM}$ . Otherwise, it will be zero. Comparison with the critical shear stress makes it possible to determine whether stick changes to slip at this point.

**PRESS**

$p$ , contact pressure at end of increment.

**DPRESS**

$\Delta p$ , increment in contact pressure.

**DDPDDH**

$\partial \Delta p / \partial \Delta h$ , current contact stiffness, in the case of soft contact (“Contact pressure-overclosure relationships,” Section 33.1.2 of the Abaqus Analysis User’s Manual).

**SLIP (NFDIR)**

Total nonrecoverable sliding motion (slip) at the beginning of the increment,  $\gamma_{\alpha}^{sl}$ . This value is the accumulated value of **DSLIP (NFDIR)** from previous increments.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Value of step time at the end of the increment.

**TIME (2)**

Value of total time at the end of the increment.

**DTIME**

Current increment in time.

**NOEL**

Element label for contact elements. Passed in as zero if contact surfaces are defined.

**CINAME**

User-specified surface interaction name associated with the friction definition, left justified. For contact elements it is the element set name given for the interface definition associated with the friction definition; if an optional name is assigned to the interface definition, **CINAME** is passed in as this name, left justified.

**SLNAME**

Slave surface name. Passed in as blank if contact elements are used.

**MSNAME**

Master surface name. Passed in as blank if contact elements are used.

**NPT**

Integration point number for contact elements. Passed in as zero if contact surfaces are defined.

## **FRIC**

### **NODE**

User-defined global slave node number (or internal node number for models defined in terms of an assembly of part instances) involved with this contact point. Corresponds to the predominant slave node of the constraint if the surface-to-surface contact formulation is used. Passed in as zero if called from a contact element.

### **NPATCH**

Not used.

### **COORDS (MCRD)**

An array containing the current coordinates of this point.

### **RCOORD (MCRD)**

If the master surface is defined as a rigid surface, this array is passed in containing the coordinates of the opposing point on the rigid surface in its current position and orientation.

### **DROT (2 , 2)**

Rotation increment matrix. For contact with a three-dimensional rigid surface, this matrix represents the incremental rotation of the surface directions relative to the rigid surface. It is provided so that vector- or tensor-valued state variables can be rotated appropriately in this subroutine. Stress and slip components are already rotated by this amount before **FRIC** is called. This matrix is passed in as a unit matrix for two-dimensional and axisymmetric contact problems.

### **TEMP (2)**

Current temperature at the slave node and the opposing master surface, respectively.

### **PREDEF (2 , NPRED)**

An array containing pairs of values of all the user-specified field variables at the end of the current increment (initial values at the beginning of the analysis and current values during the analysis). If **FRIC** is called from a contact pair, the first value in a pair corresponds to the slave node and the second value corresponds to the nearest point on the master surface. If **FRIC** is called from a large-sliding contact element, **PREDEF (1 , NPRED)** corresponds to the value at the integration point of the element and **PFREDEF (2 , NPRED)** corresponds to the nearest point on the opposing surface. If **FRIC** is called from a small-sliding contact element, **PREDEF (1 , NPRED)** corresponds to the value at the integration point of the first side and **PFREDEF (2 , NPRED)** corresponds to the value at the integration point on the opposite face of the element.

### **NFDIR**

Number of friction directions.

### **MCRD**

Number of coordinate directions at the contact point.

### **NPRED**

Number of predefined field variables.

**NSTATV**

Number of user-defined state variables.

**CHRLNGTH**

Characteristic contact surface face dimension, which can be used to define the maximum allowable elastic slip.

**PROPS (NPROPS)**

Array of user-specified property values that are used to define the frictional behavior between the contacting surfaces.

**NPROPS**

User-specified number of property values associated with this friction model.



## 1.1.9 FRIC\_COEF: User subroutine to define the frictional coefficient for contact surfaces.

**Product:** Abaqus/Standard

### References

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- “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual
- \*FRICTION
- “**FRIC\_COEF**,” Section 4.1.5 of the Abaqus Verification Manual

### Overview

---

User subroutine **FRIC\_COEF**:

- can be used to define the isotropic frictional coefficient between contacting surfaces;
- corresponds to the classical Coulomb friction model; and
- can be used with the contact pair and general contact algorithms.

### User subroutine interface

---

```

subroutine fric_coef (
C Write only -
*   fCoef, fCoefDeriv,
C Read only -
*   nBlock, nProps, nTemp, nFields,
*   jFlags, rData,
*   surfInt, surfSlv, surfMst,
*   props, slipRate, pressure,
*   tempAvg, fieldAvg )
C
    include 'aba_param.inc'
C
    dimension fCoef(nBlock),
*   fCoefDeriv(nBlock,3),
*   props(nProps),
*   slipRate(nBlock),
*   pressure(nBlock),
*   tempAvg(nBlock),
*   fieldAvg(nBlock,nFields)
C
    parameter( iKStep    = 1,

```

## FRIC\_COEF

```
*          iKInc    = 2,
*          nFlags   = 2 )
C
parameter( iTimStep = 1,
*          iTimGlb  = 2,
*          iDTimCur = 3,
*          nData    = 3 )
C
dimension jFlags(nFlags), rData(nData)
C
character*80 surfInt, surfSlv, surfMst
C
user coding to define fCoef
return
end
```

### Variables to be defined

---

#### **fCoef(nBlock)**

This array must be updated to the current values of the friction coefficient at the contact point.

#### **fCoefDeriv(nBlock,3)**

This array must be updated to the derivatives of the friction coefficient with respect to slip rate, pressure, and temperature at the contact point.

### Variables passed in for information

---

#### **nBlock**

Equal to 1.

#### **nProps**

User-specified number of property values associated with this friction model.

#### **nTemp**

1 if the temperature is defined and 0 if the temperature is not defined.

#### **nFields**

Number of user-specified field variables.

#### **jFlag(1)**

Step number.

#### **jFlag(2)**

Increment number.

**rData(1)**

Value of step time.

**rData(2)**

Value of total time.

**rData(3)**

Current increment in time from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ .

**surfInt**

User-specified surface interaction name, left justified.

**surfSlv**

Slave surface name, left justified.

**surfMst**

Master surface name, left justified.

**props(nProps)**

User-specified vector of property values to define the frictional coefficient at the contact point.

**slipRate(nBlock)**

This array contains the rate of tangential slip at the contact point for the current time increment.

**pressure(nBlock)**

This array contains the pressure at the contact point projected at the end of the current time increment.

**tempSlv(nBlock)**

Average current temperature between the master and slave surfaces at the contact point.

**fieldSlv(nFields,nBlock)**

Average current value of all the user-specified field variables between the master and slave surfaces at the contact point.



- 1.1.10 GAPCON: User subroutine to define conductance between contact surfaces or nodes in a fully coupled temperature-displacement analysis or pure heat transfer analysis.**

**Product:** Abaqus/Standard

## References

---

- “Thermal contact properties,” Section 33.2.1 of the Abaqus Analysis User’s Manual
- \*GAP CONDUCTANCE
- “**GAPCON**,” Section 4.1.6 of the Abaqus Verification Manual

## Overview

---

User subroutine **GAPCON**:

- assumes that the heat transfer between surfaces is modeled as  $q = k(\theta_A - \theta_B)$ , where  $q$  is the heat flux per unit area flowing between corresponding points  $A$  and  $B$  on the surfaces,  $k$  is the gap conductance, and  $\theta_A$  and  $\theta_B$  are the surface temperatures;
- is used to define  $k$ , providing greater flexibility than direct gap conductance definition in specifying the dependencies of  $k$  (for example, it is not necessary to define the gap conductance as a function of the average of the two surfaces’ temperatures, mass flow rates, or field variables);
- will be called at the slave nodes of a contact pair and at the integration points in a contact or a gap element for which the heat conductance definition contains a user-subroutine-defined gap conductance; and
- ignores any dependencies or data specified for the gap conductance outside the user subroutine.

## Usage with contact pairs and gap elements

---

When this subroutine is used with a contact pair, point  $A$  is on the slave surface and point  $B$  is on the master surface.

When **GAPCON** is used with gap elements of type DGAP or GAPUNIT, point  $A$  is on the first node of the element and point  $B$  is the second node of the element.

## User subroutine interface

---

```
SUBROUTINE GAPCON(AK,D,FLOWM,TEMP,PREDEF,TIME,CINAME,SLNAME,
1 MSNAME,COORDS,NOEL,NODE,NPRED,KSTEP,KINC)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CINAME,SLNAME,MSNAME
```

**C**

```
DIMENSION AK(5),D(2), FLOWM(2), TEMP(2), PREDEF(2,*),
1 TIME(2), COORDS(3)
```

*user coding to define **AK(1) -- AK(5)***

```
RETURN
END
```

## **Variables to be defined**

---

### **AK(1)**

Gap conductance,  $k$ . The units of  $k$  are energy per time (flux) per area per temperature ( $\text{JT}^{-1}\text{L}^{-2}\theta^{-1}$ ).

### **AK(2)**

$\partial k/\partial d$ , derivative of the gap conductance with respect to the clearance between the bodies. If the gap conductance is not a function of gap clearance, **AK(2)**=0.0. This variable needs to be defined only for fully coupled temperature-displacement analysis.

### **AK(3)**

$\partial k/\partial p$ , derivative of the gap conductance with respect to the pressure between the bodies. If the gap conductance is not a function of the pressure, **AK(3)**=0.0. This variable needs to be defined only for fully coupled temperature-displacement analysis.

### **AK(4)**

$\partial k/\partial \theta_A$ , derivative of the gap conductance with respect to the temperature of point  $A$  on the first surface of the interface.

### **AK(5)**

$\partial k/\partial \theta_B$ , derivative of the gap conductance with respect to the temperature of point  $B$  on the second surface of the interface.

## **Variables passed in for information**

---

### **D(1)**

Separation between the surfaces,  $d$ .

### **D(2)**

Pressure transmitted across the surfaces,  $p$ . This pressure is zero in pure heat transfer analysis.

### **FLOWM(2)**

$\dot{m}|_A, \dot{m}|_B$ , magnitudes of the mass flow rate per unit area at points  $A$  and  $B$ .

**TEMP (2)**

Current temperature at points *A* and *B*.

**PREDEF (2 , NPRED)**

An array containing pairs of values of all of the user-specified field variables at the end of the current increment at points *A* and *B* (initial values at the beginning of the analysis and current values during the analysis).

**TIME (1)**

Value of step time at the end of the increment.

**TIME (2)**

Value of total time at the end of the increment.

**CINAME**

User-specified surface interaction name associated with the heat conductance definition, left justified. For contact elements it is the element set name given for the interface definition associated with the heat conductance definition; if an optional name is assigned to the interface definition, **CINAME** is passed in as this name, left justified. For gap elements it is the element set name for the element definition associated with the heat conductance definition.

**SLNAME**

Slave surface name. Passed in as blank if contact or gap elements are used.

**MSNAME**

Master surface name. Passed in as blank if contact or gap elements are used.

**COORDS**

An array containing the coordinates of point *A*. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**NOEL**

Element label for contact or gap elements. Passed in as zero if contact surfaces are defined.

**NODE**

Slave node number (point *A*) if **GAPCON** is called for a contact pair.

**NPRED**

Number of predefined field variables.

**KSTEP**

Step number.

**KINC**

Increment number.

### 1.1.11     **GAPELECTR: User subroutine to define electrical conductance between surfaces in a coupled thermal-electrical analysis.**

**Product:** Abaqus/Standard

#### References

---

- “Electrical contact properties,” Section 33.3.1 of the Abaqus Analysis User’s Manual
- \*GAP ELECTRICAL CONDUCTANCE

#### Overview

---

User subroutine **GAPELECTR**:

- assumes that the electrical current flowing between the interface surfaces is modeled as  $J = \sigma_g(\varphi_A - \varphi_B)$ , where  $J$  is the electrical current density flowing across the interface from point  $A$  (the slave surface) to point  $B$  (the master surface),  $\varphi_A$  and  $\varphi_B$  are the electrical potential on opposite points of the surfaces, and  $\sigma_g$  is the surface electrical conductance;
- is used to define  $\sigma_g$ , providing much greater flexibility than direct gap electrical conductance definition in specifying the dependencies of  $\sigma_g$  (for instance, it is not necessary to define the gap electrical conductance as a function of the average of the two surfaces’ temperatures and/or field variables);
- will be called at the slave nodes of a contact pair (“Defining contact pairs in Abaqus/Standard,” Section 32.3.1 of the Abaqus Analysis User’s Manual) for which the gap electrical conductance is defined in a user subroutine; and
- ignores any dependencies or data specified for the gap electrical conductance outside the user subroutine.

#### User subroutine interface

---

```

SUBROUTINE GAPELECTR(SIGMA,D,TEMP,PREDEF,TIME,CINAME,
1 SLNAME,MSNAME,COORDS,NODE,NPRED,KSTEP,KINC)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CINAME,SLNAME,MSNAME
C
DIMENSION SIGMA(3),TEMP(2),PREDEF(2,*),TIME(2),COORDS(2,3)

user coding to define SIGMA(1) -- SIGMA(3)

RETURN
END

```

**Variables to be defined**

---

**SIGMA (1)**

Gap electrical conductance,  $\sigma_g$ .

**SIGMA (2)**

$\partial\sigma_g/\partial\theta_A$ , derivative of the gap electrical conductance with respect to the temperature of point *A*. If the gap electrical conductance is not a function of  $\theta_A$ , **SIGMA (2)** = 0.0.

**SIGMA (3)**

$\partial\sigma_g/\partial\theta_B$ , derivative of the gap electrical conductance with respect to the temperature of point *B*. If the gap electrical conductance is not a function of  $\theta_B$ , **SIGMA (3)** = 0.0.

**Variables passed in for information**

---

**D**

Separation between the interface surfaces, *d*.

**TEMP (2)**

Current temperature at points *A* and *B*.

**PREDEF (2 ,NPRED)**

An array containing pairs of values of all of the user-specified field variables at the end of the current increment at points *A* and *B* (initial values at the beginning of the analysis and current values during the analysis).

**TIME (1)**

Value of step time at the end of the increment.

**TIME (2)**

Value of total time at the end of the increment.

**CINAME**

User-specified surface interaction name, left justified.

**SLNAME**

Slave surface name.

**MSNAME**

Master surface name.

**COORDS**

An array containing the current coordinates of points *A* and *B*. **COORDS (1 ,K1)** are the coordinates at point *A*, and **COORDS (2 ,K1)** are the coordinates at point *B*.

**NODE**

Slave node number (point *A*).

**NPRED**

Number of predefined field variables.

**KSTEP**

Step number.

**KINC**

Increment number.



### 1.1.12 HARDINI: User subroutine to define initial equivalent plastic strain and initial backstress tensor.

**Product:** Abaqus/Standard

#### References

---

- “Initial conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.2.1 of the Abaqus Analysis User’s Manual
- “Classical metal plasticity,” Section 20.2.1 of the Abaqus Analysis User’s Manual
- “Models for metals subjected to cyclic loading,” Section 20.2.2 of the Abaqus Analysis User’s Manual
- “Extended Drucker-Prager models,” Section 20.3.1 of the Abaqus Analysis User’s Manual
- \*INITIAL CONDITIONS
- “**HARDINI**,” Section 4.1.8 of the Abaqus Verification Manual

#### Overview

---

User subroutine **HARDINI**:

- can be used only for material models that use metal plasticity or Drucker-Prager plasticity;
- can be used to provide initial equivalent plastic strain values as a function of element number, material point number, and/or material point coordinates for isotropic and combined hardening;
- enables you to specify initial conditions for the backstress tensor as a function of element number, material point number, and/or material point coordinates for kinematic and combined hardening;
- will be called to define the initial equivalent plastic strain and, if relevant, the initial backstresses at material points for which user-subroutine-defined initial hardening conditions are specified; and
- is intended for use when the initial equivalent plastic strain and/or backstress distributions are too complicated to specify directly as initial hardening conditions.

#### Defining backstress components

---

The number of backstress components that must be defined depends on the element type for which this routine is being called. Part VI, “Elements,” of the Abaqus Analysis User’s Manual describes the number of stress components for each element type; the number of backstress components is identical to the number of stress components. The order of the backstress components is the same as the order of the stress components. For example, in three-dimensional continuum elements six backstress components must be defined in the order  $\alpha_{11}, \alpha_{22}, \alpha_{33}, \alpha_{12}, \alpha_{13}, \alpha_{23}$ .

## User subroutine interface

---

```
SUBROUTINE HARDINI (ALPHA,EQPS,COORDS,NTENS,NCRDS,NOEL,NPT,
1 LAYER,KSPT,LREBAR,REBARN)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION ALPHA (NTENS,*), COORDS (NCRDS)
CHARACTER*80 REBARN
```

*user coding to define EQPS and, if relevant, ALPHA (NTENS)*

```
RETURN
END
```

## Variables to be defined

---

The variables described below are element-type dependent.

### **EQPS**

Equivalent plastic strain.

### **ALPHA (1,1)**

First backstress component of the first backstress.

### **ALPHA (2,1)**

Second backstress component of the first backstress.

### **ALPHA (3,1)**

Third backstress component of the first backstress.

### **Etc.**

**NTENS** backstress component values should be defined for each backstress.

## Variables passed in for information

---

### **COORDS**

An array containing the initial coordinates of this point.

### **NTENS**

Number of backstress values to be defined. This number depends on the element type.

**NCRDS**

Number of coordinates.

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**LREBAR**

Rebar flag. If **LREBAR**=1, the current integration point is associated with element rebar. Otherwise, **LREBAR**=0.

**REBARN**

Name of the rebar to which the current integration point belongs, which is the name given in the rebar or rebar layer definition (“Defining reinforcement,” Section 2.2.3 of the Abaqus Analysis User’s Manual, or “Defining rebar as an element property,” Section 2.2.4 of the Abaqus Analysis User’s Manual). If no name was given in the rebar or rebar layer definition, this variable will be blank. This variable is relevant only when **LREBAR**=1.



**1.1.13 HETVAL: User subroutine to provide internal heat generation in heat transfer analysis.**

**Product:** Abaqus/Standard

## References

---

- “Uncoupled heat transfer analysis,” Section 6.5.2 of the Abaqus Analysis User’s Manual
- “Fully coupled thermal-stress analysis,” Section 6.5.4 of the Abaqus Analysis User’s Manual
- \*HEAT GENERATION
- “**HETVAL**,” Section 4.1.9 of the Abaqus Verification Manual

## Overview

---

User subroutine **HETVAL**:

- can be used to define a heat flux due to internal heat generation in a material, for example, as might be associated with phase changes occurring during the solution;
- allows for the dependence of internal heat generation on state variables (such as the fraction of material transformed) that themselves evolve with the solution and are stored as solution-dependent state variables;
- will be called at all material calculation points for which the material definition contains volumetric heat generation during heat transfer, coupled temperature-displacement, or coupled thermal-electrical analysis procedures;
- can be useful if it is necessary to include a kinetic theory for a phase change associated with latent heat release (for example, in the prediction of crystallization in a polymer casting process);
- can be used in conjunction with user subroutine **USDFLD** if it is desired to redefine any field variables before they are passed in; and
- cannot be used with user subroutine **UMATHT**.

## User subroutine interface

---

```

SUBROUTINE HETVAL(CMNAME, TEMP, TIME, DTIME, STATEV, FLUX,
1 PREDEF, DPRED)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
C
DIMENSION TEMP(2), STATEV(*), PREDEF(*), TIME(2), FLUX(2),
1 DPRED(*)

```

## HETVAL

*user coding to define **FLUX** and update **STATEV***

```
RETURN  
END
```

### Variables to be defined

---

#### **FLUX (1)**

Heat flux,  $r$  (thermal energy per time per volume:  $JT^{-1}L^{-3}$ ), at this material calculation point.

#### **FLUX (2)**

Rate of change of heat flux per temperature,  $\partial r / \partial \theta$ . This variable is nonzero only if the heat flux depends on temperature. It is needed to define a correct Jacobian matrix.

### Variable that can be updated

---

#### **STATEV (\*)**

An array containing the user-defined solution-dependent state variables at this point.

In an uncoupled heat transfer analysis **STATEV** is passed into subroutine **HETVAL** as the values of these variables at the beginning of the increment. However, any updating of **STATEV** in user subroutine **USDFLD** will be included in the values passed into subroutine **HETVAL** since this routine is called before **HETVAL**. In addition, if **HETVAL** is being used in a fully coupled temperature-displacement analysis and user subroutine **UEXPAN**, user subroutine **CREEP**, user subroutine **UMAT**, or user subroutine **UTRS** is used to define the mechanical behavior of the material, those routines are called before this routine; therefore, any updating of **STATEV** done in **UEXPAN**, **CREEP**, **UMAT**, or **UTRS** will be included in the values passed into this routine.

In all cases **STATEV** should be passed back from user subroutine **HETVAL** containing the values of the state variables at the end of the current increment.

### Variables passed in for information

---

#### **CMNAME**

User-specified material name, left justified.

#### **TEMP (1)**

Current temperature.

#### **TEMP (2)**

Temperature increment.

#### **TIME (1)**

Step time at the end of the increment.

**TIME (2)**

Total time at the end of the increment.

**DTIME**

Time increment.

**PREDEF (\*)**

An array containing the values of all of the user-specified field variables at this point (initial values at the beginning of the analysis and current values during the analysis).

**DPRED (\*)**

Array of increments of predefined field variables.



### 1.1.14 MPC: User subroutine to define multi-point constraints.

**Product:** Abaqus/Standard

#### References

---

- “General multi-point constraints,” Section 31.2.2 of the Abaqus Analysis User’s Manual
- \*MPC

#### Overview

---

User subroutine **MPC**:

- is called to impose a user-defined multi-point constraint and is intended for use when general constraints cannot be defined with one of the MPC types provided by Abaqus/Standard;
- can use only degrees of freedom that also exist on an element somewhere in the same model (methods for overcoming this limitation are discussed below);
- can generate linear as well as nonlinear constraints;
- allows definition of constraints involving finite rotations; and
- makes it possible to switch constraints on and off during an analysis.

#### Coding methods

---

There are two methods for coding this routine. By default, the subroutine operates in a *degree of freedom* mode. In this mode each call to this subroutine allows one individual degree of freedom to be constrained. Alternatively, you can specify that the subroutine operate in a *nodal* mode. In this mode each call to this subroutine allows a set of constraints to be imposed all at once; that is, on multiple degrees of freedom of the dependent node. In either case, the routine will be called for each user-subroutine-defined multi-point constraint or set of constraints. See “General multi-point constraints,” Section 31.2.2 of the Abaqus Analysis User’s Manual, for details.

#### Constraints that involve rotational degrees of freedom

In geometrically nonlinear analyses Abaqus/Standard compounds three-dimensional rotations based on a finite-rotation formulation and not by simple addition of the individual rotation components (see “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual, and “Rotation variables,” Section 1.3.1 of the Abaqus Theory Manual). An incremental rotation involving one component usually results in changes in all three total rotation components. Therefore, any general constraint that involves large three-dimensional rotations should be implemented using the nodal mode of user subroutine **MPC**. The single degree of freedom version of user subroutine **MPC** can be used for geometrically linear problems, geometrically nonlinear problems with planar rotations, and constraints that do not involve rotation components.

## Constraints that involve degrees of freedom that are not active in the model

---

The degrees of freedom involved in user MPCs must appear on some element or Abaqus/Standard MPC type in the model: user MPCs cannot use degrees of freedom that have not been introduced somewhere on an element. For example, a mesh that uses only continuum (solid) elements cannot have user MPCs that involve rotational degrees of freedom. The simplest way to overcome this limitation is to introduce an element somewhere in the model that uses the required degrees of freedom but does not affect the solution in any other way. Alternatively, if the degrees of freedom are rotations, they can be activated by the use of a library BEAM-type MPC somewhere in the model.

## Use with nodal coordinate systems

---

When a local coordinate system (“Transformed coordinate systems,” Section 2.1.5 of the Abaqus Analysis User’s Manual) and a user MPC are both used at a node, the variables at the node are first transformed before the MPC is imposed. Therefore, user-supplied MPCs must be based on the transformed degrees of freedom. The local-to-global transformation matrices  $\mathbf{T}^I$  for the individual nodes:

$$\mathbf{u}_{global}^I = \mathbf{T}^I \cdot \mathbf{u}_{local}^I$$

are passed in for information.

## Degree of freedom version of user subroutine MPC

---

This version of user subroutine **MPC** allows for one individual degree of freedom to be constrained and, thus, eliminated at a time. The constraint can be quite general and nonlinear of the form:

$$f(u^1, u^2, u^3, \dots, u^N, \text{geometry, temperature, field variables}) = 0.$$

The first degree of freedom in this function,  $u^1$ , is the degree of freedom that will be eliminated to impose the constraint.  $u^2$ ,  $u^3$ , etc. are any other degrees of freedom that are involved in the constraint. Since  $u^1$  will be eliminated to impose the constraint, it cannot be used in subsequent kinematic constraints (multi-point constraints, linear equation constraints, or boundary conditions). Therefore, the user MPCs are imposed in the order given in the input.

You must provide, at all times, two items of information in user subroutine **MPC**:

1. A list of degree of freedom identifiers at the nodes that are listed in the corresponding multi-point constraint definition. This list corresponds to  $u^1, u^2, u^3$ , etc. in the constraint as given above.
2. An array of the derivatives

$$A^1 = \frac{\partial f}{\partial u^1}, \quad A^2 = \frac{\partial f}{\partial u^2}, \quad A^3 = \frac{\partial f}{\partial u^3}, \quad \dots$$

of the constraint function with respect to the degrees of freedom involved. This array is needed for the redistribution of loads from degree of freedom  $u^1$  to the other degrees of freedom and for the elimination of  $u^1$  from the system matrices.

In addition, you can provide the value of the dependent degree of freedom  $u^1$  as a function of the independent degrees of freedom  $u^2, u^3$ , etc. If this value is not provided, Abaqus/Standard will update  $u^1$  based on the linearized form of the constraint equation as

$$u^1 = -\frac{1}{A^1} \sum_{i=2}^N A^i u^i.$$

Subroutine **MPC** should be coded and checked with care: if the array of derivatives  $\partial f / \partial u^1$ , etc. does not correspond to the definition of  $u^1$  in terms of  $u^2, u^3$ , etc., forces will be transmitted improperly by the MPC and violations of equilibrium may occur. In addition, convergence of the solution may be adversely affected.

## User subroutine interface

---

```

SUBROUTINE MPC(UE,A,JDOF,MDOF,N,JTYPE,X,U,UINIT,MAXDOF,
* LMPC,KSTEP,KINC,TIME,NT,NF,TEMP,FIELD,LTRAN,TRAN)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION A(N),JDOF(N),X(6,N),U(MAXDOF,N),UINIT(MAXDOF,N),
* TIME(2),TEMP(NT,N),FIELD(NF,NT,N),LTRAN(N),TRAN(3,3,N)

```

*user coding to define UE, A, JDOF, and, optionally, LMPC*

```

RETURN
END

```

## Variables to be defined

---

### A (N)

An array containing the derivatives of the constraint function,

$$A(1) = \frac{\partial f}{\partial u^1}, \quad A(2) = \frac{\partial f}{\partial u^2}, \quad \dots$$

The coding in the subroutine must define **N** entries in **A**, where **N** is defined below.

**JDOF (N)**

An array containing the degree of freedom identifiers at the nodes that are involved in the constraint. For example, if  $u^1$  is the  $z$ -displacement at a node, give **JDOF**(1) = 3; if  $u^2$  is the  $x$ -displacement at a node, give **JDOF**(2) = 1. The coding in the subroutine must define **N** entries in **JDOF**, where **N** is defined below.

**Variables that can be updated**

---

**UE**

This variable is passed in as the total value of the eliminated degree of freedom,  $u^1$ . This variable will either be zero or have the current value of  $u^1$  based on the linearized constraint equation, depending at which stage of the iteration the user subroutine is called. If the constraint is linear and is used in a small-displacement analysis (nonlinear geometric effects are not considered) or in a perturbation analysis, this variable need not be defined: Abaqus/Standard will compute  $u^1$  as

$$u^1 = -\frac{1}{A^1} \sum_{i=2}^N A^i u^i.$$

If the constraint is nonlinear, this variable should be updated to the value of  $u^1$  at the end of the increment to satisfy the constraint exactly. If the return value is the same as the incoming value, Abaqus/Standard will update the eliminated degree of freedom based on the linearized form of the constraint equation. In this case the constraint is not likely to be satisfied exactly.

**LMPC**

Set this variable to zero to avoid the application of the multi-point constraint. The MPC will be applied if the variable is not changed. This variable must be set to zero every time the subroutine is called if the user MPC is to remain deactivated. This MPC variable is useful for switching the MPC on and off during an analysis. However, the option should be used with care: switching off an MPC may cause a sudden disturbance in equilibrium, which can lead to convergence problems. If this variable is used to switch on an MPC during an analysis, the variable **UE** should be defined; otherwise, the constraint may not be satisfied properly.

**Variables passed in for information**

---

**MDOF**

Maximum number of active degrees of freedom per node involved in the MPC. For the degree of freedom mode of user subroutine **MPC**, **MDOF**= 1.

**N**

Number of degrees of freedom that are involved in the constraint, defined as the number of nodes given in the corresponding multi-point constraint definition. If more than one degree of freedom at a node is

involved in a constraint, the node must be repeated as needed or, alternatively, the nodal mode should be used.

#### **JTYPE**

Constraint identifier given for the corresponding multi-point constraint definition.

#### **X (6,N)**

An array containing the original coordinates of the nodes involved in the constraint.

#### **U (MAXDOF , N)**

An array containing the values of the degrees of freedom at the nodes involved in the constraint. These values will be either the values at the end of the previous iteration or the current values based on the linearized constraint equation, depending at which stage of the iteration the user subroutine is called.

#### **UINIT (MAXDOF , N)**

An array containing the values at the beginning of the current iteration of the degrees of freedom at the nodes involved in the constraint. This information is useful for decision-making purposes when you do not want the outcome of a decision to change during the course of an iteration. For example, there are constraints in which the degree of freedom to be eliminated changes during the course of the analysis, but it is necessary to prevent the choice of the dependent degree of freedom from changing during the course of an iteration.

#### **MAXDOF**

Maximum degree of freedom number at any node in the analysis. For example, for a coupled temperature-displacement analysis with continuum elements, **MAXDOF** will be equal to 11.

#### **KSTEP**

Step number.

#### **KINC**

Increment number within the step.

#### **TIME (1)**

Current value of step time.

#### **TIME (2)**

Current value of total time.

#### **NT**

Number of positions through a section where temperature or field variable values are stored at a node. In a mesh containing only continuum elements, **NT**=1. For a mesh containing shell or beam elements, **NT** is the largest of the values specified for the number of temperature points in the shell or beam section definition (or 2 for temperatures specified together with gradients for shells or two-dimensional beams, 3 for temperatures specified together with gradients for three-dimensional beams).

**NF**

Number of different predefined field variables requested for any node (including field variables defined as initial conditions).

**TEMP (NT , N)**

An array containing the temperatures at the nodes involved in the constraint. This array is not used for a heat transfer, coupled temperature-displacement, or coupled thermal-electrical analysis since the temperatures are degrees of freedom of the problem.

**FIELD (NF , NT , N)**

An array containing all field variables at the nodes involved in the constraint.

**LTRAN (N)**

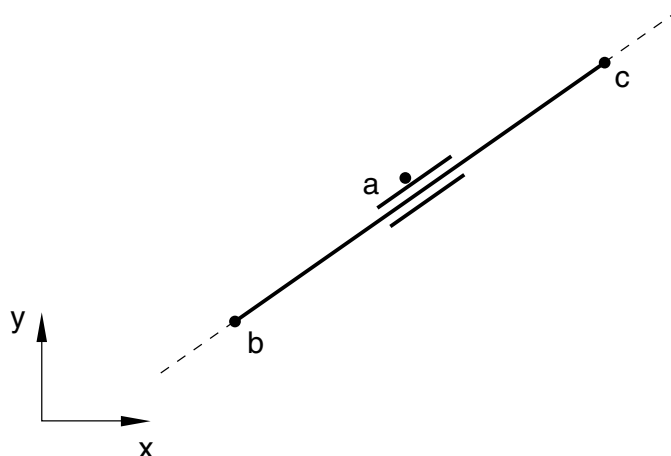
An integer array indicating whether the nodes in the MPC are transformed. If **LTRAN (I)=1**, a transformation is applied to node **I**; if **LTRAN (I)=0**, no transformation is applied.

**TRAN (3 , 3 , N)**

An array containing the local-to-global transformation matrices for the nodes used in the MPC. If no transformation is present at node **I**, **TRAN (\* , \* , I)** is the identity matrix.

**Example: Nonlinear single degree of freedom MPC**

An example of a nonlinear single degree of freedom MPC is a geometrically nonlinear two-dimensional slider involving nodes *a*, *b*, and *c*. The constraint forces node *a* to be on the straight line connecting nodes *b* and *c* (see Figure 1.1.14–1).



**Figure 1.1.14–1** Nonlinear MPC example: two-dimensional slider.

The constraint equation can be written in the form

$$f(u^a, v^a, u^b, v^b, u^c, v^c) = (x^a - x^b)(y^c - y^b) - (y^a - y^b)(x^c - x^b) = 0,$$

where  $(x^a, y^a)$ ,  $(x^b, y^b)$ , and  $(x^c, y^c)$  are the current locations of  $a$ ,  $b$ , and  $c$ . The derivatives are readily obtained as

$$\frac{\partial f}{\partial u^a} = y^c - y^b, \quad \frac{\partial f}{\partial u^b} = y^a - y^c, \quad \frac{\partial f}{\partial u^c} = y^b - y^a,$$

$$\frac{\partial f}{\partial v^a} = x^b - x^c, \quad \frac{\partial f}{\partial v^b} = x^c - x^a, \quad \frac{\partial f}{\partial v^c} = x^a - x^b.$$

Depending on the orientation of the segment  $(b, c)$ , we choose either  $u^a$  or  $v^a$  as the degree of freedom to be eliminated. If  $|x^c - x^b| \geq |y^c - y^b|$ , we choose  $v^a$  as the dependent degree of freedom. If  $|x^c - x^b| < |y^c - y^b|$ , we choose  $u^a$  as the dependent degree of freedom. Moreover, if points  $b$  and  $c$  are coincident, the constraint is not applied.

To prevent the choice of either  $u^a$  or  $v^a$  as the dependent degree of freedom from changing during the course of an iteration, the orientation of the segment  $(b, c)$  is tested based on the geometry at the beginning of the iteration. The dependent degree of freedom is allowed to change from increment to increment.

Suppose the above multi-point constraint is defined as type 1, with nodes  $a$ ,  $a$ ,  $b$ ,  $b$ ,  $c$ ,  $c$ . The user subroutine **MPC** could be coded as follows:

```

SUBROUTINE MPC(UE,A,JDOF,MDOF,N,JTYPE,X,U,UINIT,MAXDOF,
* LMPC,KSTEP,KINC,TIME,NT,NF,TEMP,FIELD,LTRAN,TRAN)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION A(N),JDOF(N),X(6,N),U(MAXDOF,N),UINIT(MAXDOF,N),
* TIME(2),TEMP(NT,N),FIELD(NF,NT,N),LTRAN(N),TRAN(3,3,N)
PARAMETER( PRECIS = 1.D-15 )
C
IF (JTYPE .EQ. 1) THEN
  DYBC0 = X(2,5) + UINIT(2,5) - X(2,3) - UINIT(2,3)
  DXBC0 = X(1,3) + UINIT(1,3) - X(1,5) - UINIT(1,5)
  DYBC = X(2,5) + U(2,5) - X(2,3) - U(2,3)
  DXBC = X(1,3) + U(1,3) - X(1,5) - U(1,5)
  A(3) = X(2,1) + U(2,1) - X(2,5) - U(2,5)
  A(4) = X(1,5) + U(1,5) - X(1,1) - U(1,1)
  A(5) = X(2,3) + U(2,3) - X(2,1) - U(2,1)
  A(6) = X(1,1) + U(1,1) - X(1,3) - U(1,3)
  JDOF(3) = 1

```

```

JDOF(4) = 2
JDOF(5) = 1
JDOF(6) = 2
IF (ABS(DYBC0).LE.PRECIS .AND. ABS(DXBC0).LE.PRECIS) THEN
C
C      POINTS B AND C HAVE COLLAPSED. DO NOT APPLY CONSTRAINT.
C
C      LMPC = 0
ELSE IF (ABS(DXBC0) .LT. ABS(DYBC0)) THEN
C
C      MAKE U_A DEPENDENT DOF.
C
C      JDOF(1) = 1
C      JDOF(2) = 2
C      A(1) = DYBC
C      A(2) = DXBC
C      UE = A(5)A(2)/A(1) + X(1,3) + U(1,3) - X(1,1)
ELSE
C
C      MAKE V_A DEPENDENT DOF.
C
C      JDOF(1) = 2
C      JDOF(2) = 1
C      A(1) = DXBC
C      A(2) = DYBC
C      UE = -A(6)A(2)/A(1) + X(2,3) + U(2,3) - X(2,1)
END IF
END IF
C
RETURN
END

```

### Nodal version of user subroutine MPC

---

The nodal version of user subroutine **MPC** allows for multiple degrees of freedom of a node to be eliminated simultaneously. The set of constraints can be quite general and nonlinear, of the form

$$f_i(u^1, u^2, u^3, \dots, u^N, \text{geometry, temperature, field variables}) = 0 \quad i = 1, 2, \dots, \text{NDEP}.$$

**NDEP** is the number of dependent degrees of freedom that are involved in the constraint and should have a value between 1 and **MDOF**, which is the number of active degrees of freedom per node in the analysis. *N* is the number of nodes involved in the constraint. The scalar constraint functions  $f_i$  can also

be considered as a vector function  $f$ , and the first set of degrees of freedom  $u^1$  in the vector function  $f$  will be eliminated to impose the constraint. The sets  $u^2$ ,  $u^3$ , etc. are the independent degrees of freedom at nodes 2, 3, etc. involved in the constraint. The set  $u^1$  must be composed of **NDEP** degrees of freedom at the first node of the MPC definition. For example, if the dependent degrees of freedom are the  $x$ -displacement, the  $z$ -displacement, and the  $y$ -rotation at the first node,  $u^1 = (u_x^1, u_z^1, \phi_y^1)$ .  $u^2$ ,  $u^3$ , etc. can be composed of any number of degrees of freedom, depending on which ones play a role in the constraint, and need not be of the same size; for example,  $u^2 = (u_y^2)$  and  $u^3 = (u_x^3, u_y^3, u_z^3)$ .

The dependent node can also reappear as an independent node in the MPC. However, since the dependent degrees of freedom of this node will be eliminated, they cannot be used as independent degrees of freedom in this MPC. For example, if the rotations at node  $a$  are constrained by the MPC, the displacements of node  $a$  can still be used as independent degrees of freedom in the MPC, but the rotations themselves cannot. Similarly, the degrees of freedom that will be eliminated to impose the constraint cannot be used in subsequent kinematic constraints (multi-point constraints, linear equation constraints, or boundary conditions). The MPCs are imposed in the order given in the input for this purpose.

The nodal version of user subroutine **MPC** was designed with the application of nonlinear constraints involving large three-dimensional rotations in mind. Due to the incremental nature of the solution procedure in Abaqus/Standard, a linearized set of constraints

$$\delta f_i = A_i^1 \cdot \delta u^1 + A_i^2 \cdot \delta u^2 + A_i^3 \cdot \delta u^3 + \dots = 0 \quad i = 1, 2, \dots, \text{NDEP},$$

where  $A_i^1 = A_i^1(u^1, u^2, \dots)$ ,  $A_i^2 = A_i^2(u^1, u^2, \dots)$ , etc. is applied during each iteration. This linearized set of constraints is used for the calculation of equilibrium. For finite rotations the linearized equation is given in terms of the linearized rotations  $\delta\theta^1, \delta\theta^2, \delta\theta^3, \dots$ , yielding

$$\delta f_i = A_i^1 \cdot \delta\theta^1 + A_i^2 \cdot \delta\theta^2 + A_i^3 \cdot \delta\theta^3 + \dots = 0 \quad i = 1, 2, \dots, \text{NDEP}.$$

Since the linearized rotation field,  $\delta\theta$ , is not the variation of the total rotation vector,  $\phi$  (see “Rotation variables,” Section 1.3.1 of the Abaqus Theory Manual), you cannot obtain the linearized constraint equation by simply taking derivatives of the vector function,  $f$ , with respect to the rotational degrees of freedom involved. The formulation of the linearized constraint in  $\delta\theta$  is equivalent to the formulation of a geometrically linear constraint in the deformed configuration and is generally easier to formulate than the constraint in terms of  $\delta\phi$ . For an exact formulation of the constraint, the dependent components of the total rotation vector  $\phi^1$  must be defined exactly (see “Rotation variables,” Section 1.3.1 of the Abaqus Theory Manual).

You must provide, at all times, two items of information in subroutine **MPC**:

1. A matrix of degree of freedom identifiers at the nodes that are listed in the corresponding multi-point constraint definition. The columns of this matrix correspond to  $u^1, u^2, u^3$ , etc. in the set of constraints as given above, where unused entries are padded with zeros. The number of nonzero entries in  $u^1$  will implicitly determine the number of dependent degrees of freedom, **NDEP**.
2. The matrices representing the linearized constraint function with respect to the degrees of freedom involved. These matrices are needed for the redistribution of loads from degrees of freedom  $u^1$  to

the other degrees of freedom and for the elimination of  $u^1$  from the system matrices. For constraints that do not involve three-dimensional rotations and constraints with planar rotations, these matrices can be readily obtained from the derivatives of the total constraint function with respect to the degrees of freedom involved:

$$A_{ij}^1 = \frac{\partial f_i}{\partial u_j^1}, \quad A_{ij}^2 = \frac{\partial f_i}{\partial u_j^2}, \quad A_{ij}^3 = \frac{\partial f_i}{\partial u_j^3}, \quad \dots$$

For constraints that involve finite rotations, the matrices follow from the linearized form:

$$A_{ij}^1 \delta\theta_j^1 + A_{ij}^2 \delta\theta_j^2 + A_{ij}^3 \delta\theta_j^3 + \dots = 0.$$

In addition, you can provide the values of the dependent degrees of freedom  $u^1$ , as a function of the independent degrees of freedom  $u^2, u^3$ , etc. For finite rotations,  $\phi^1$  must be specified as a function of  $\phi^2, \phi^3$ , etc. If these values are not provided, Abaqus/Standard will update  $u^1$  based on the linearized form of the constraint equations. Subroutine **MPC** should be coded and checked with care: if the matrices of derivatives  $A_{ij}^1$ , etc. do not correspond to the definition of  $u^1$  in terms of  $u^2, u^3$ , etc., forces will be transmitted improperly by the MPC and violations of equilibrium may occur. In addition, convergence of the solution may be adversely affected.

## User subroutine interface

---

```

SUBROUTINE MPC (UE ,A ,JDOF ,MDOF ,N ,JTYPE ,X ,U ,UINIT ,MAXDOF ,
* LMPC ,KSTEP ,KINC ,TIME ,NT ,NF ,TEMP ,FIELD ,LTRAN ,TRAN)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION UE (MDOF) ,A (MDOF,MDOF,N) ,JDOF (MDOF,N) ,X (6,N) ,
* U (MAXDOF,N) ,UINIT (MAXDOF,N) ,TIME (2) ,TEMP (NT,N) ,
* FIELD (NF,NT,N) ,LTRAN (N) ,TRAN (3,3,N)

```

*user coding to define JDOF, UE, A and, optionally, LMPC*

```

RETURN
END

```

## Variables to be defined

---

### **JDOF (MDOF , N)**

Matrix of degrees of freedom identifiers at the nodes involved in the constraint. Before each call to **MPC**, Abaqus/Standard will initialize all of the entries of **JDOF** to zero. All active degrees of freedom for a given column (first index ranging from 1 to **MDOF**) must be defined starting at the top of the column with no zeros in between. A zero will mark the end of the list for that column. The number of nonzero entries in the first column will implicitly determine the number of dependent degrees of freedom (**NDEP**). For example, if the dependent degrees of freedom are the *z*-displacement, the *x*-rotation, and the *z*-rotation at the first node, **NDEP** = 3 and

$$\text{JDOF}(1, 1) = 3, \quad \text{JDOF}(2, 1) = 4, \quad \text{JDOF}(3, 1) = 6.$$

If the degrees of freedom at the third node involved in the MPC are the *x*-displacement and the *y*-rotation, define

$$\text{JDOF}(1, 3) = 1, \quad \text{JDOF}(2, 3) = 5.$$

### **A (MDOF , MDOF , N)**

Submatrices of coefficients of the linearized constraint function,

$$\mathbf{A}(\mathbf{I}, \mathbf{J}, 1) = A_{ij}^1, \quad \mathbf{A}(\mathbf{I}, \mathbf{J}, 2) = A_{ij}^2, \quad \dots$$

Before each call to user subroutine **MPC**, Abaqus/Standard will initialize all of the entries of **A** to zero; therefore, only nonzero entries need to be defined. If the coding in the subroutine defines **NDEP** nonzero entries in the column **JDOF (J , 1)**, it should define **NDEP** × **NDEP** entries in the submatrix **A (I , J , 1)**. Since this submatrix will be inverted to impose the MPC, it must be nonsingular. A maximum of **NDEP** × **MDOF** entries can be defined for the remaining submatrices **A (I , J , K)**, **K** = 2, …, **N**. The number of columns in each submatrix **A (I , J , K)** must correspond to the number of nonzero entries in the corresponding column of the matrix **JDOF (J , K)**.

## Variables that can be updated

---

### **UE (NDEP)**

This array is passed in as the total value of the eliminated degrees of freedom,  $\mathbf{u}^1$ . This array will either be zero or contain the current values of  $\mathbf{u}^1$  based on the linearized constraint equations, depending at which stage of the iteration the user subroutine is called. For small-displacement analysis or perturbation analysis this array need not be defined: Abaqus/Standard will compute  $\mathbf{u}^1$  as

$$u_i^1 = - \sum_{r=1}^{\text{NDEP}} A_{ir}^1 {}^{-1} \sum_{s=2}^{\text{N}} \sum_{t=1}^{\text{MDOF}} A_{rt}^s u_{\text{JDOF}(t,s)}^s \quad i = 1, \dots, \text{NDEP}.$$

For large-displacement analysis this array can be updated to the value of  $u^1$  at the end of the increment to satisfy the constraint exactly. If the return values are the same as the incoming values, Abaqus/Standard will update the eliminated degrees of freedom based on the linearized form of the constraint equations. In this case the constraint is not likely to be satisfied exactly.

**LMPC**

Set this variable to zero to avoid the application of the multi-point constraint. If the variable is not changed, the MPC will be applied. This variable must be set to zero every time the subroutine is called if the user MPC is to remain deactivated. This MPC variable is useful for switching the MPC on and off during an analysis. This option should be used with care: switching off an MPC may cause a sudden disturbance in equilibrium, which can lead to convergence problems.

---

**Variables passed in for information**

---

**MDOF**

Number of active degrees of freedom per node in the analysis. For example, for a coupled temperature-displacement analysis with two-dimensional continuum elements, the active degrees of freedom are 1, 2, and 11 and, hence, **MDOF** will be equal to 3.

**N**

Number of nodes involved in the constraint. The value of **N** is defined as the number of nodes given in the corresponding multi-point constraint definition.

**JTYPE**

Constraint identifier given for the corresponding multi-point constraint definition.

**X(6,N)**

An array containing the original coordinates of the nodes involved in the constraint.

**U(MAXDOF,N)**

An array containing the values of the degrees of freedom at the nodes involved in the constraint. These values will either be the values at the end of the previous iteration or the current values based on the linearized constraint equation, depending at which stage of the iteration the user subroutine is called.

**UINIT (MAXDOF,N)**

An array containing the values at the beginning of the current iteration of the degrees of freedom at the nodes involved in the constraint. This information is useful for decision-making purposes when you do not want the outcome of a decision to change during the course of an iteration. For example, there are constraints in which the degrees of freedom to be eliminated change during the course of the analysis, but it is necessary to prevent the choice of the dependent degrees of freedom from changing during the course of an iteration.

**MAXDOF**

Maximum degree of freedom number at any node in the analysis. For example, for a coupled temperature-displacement analysis with continuum elements, **MAXDOF** is equal to 11.

**KSTEP**

Step number.

**KINC**

Increment number within the step.

**TIME (1)**

Current value of step time.

**TIME (2)**

Current value of total time.

**NT**

Number of positions through a section where temperature or field variable values are stored at a node. In a mesh containing only continuum elements, **NT**=1. For a mesh containing shell or beam elements, **NT** is the largest of the values specified for the number of temperature points in the shell or beam section definition (or 2 for temperatures specified together with gradients for shells or two-dimensional beams, 3 for temperatures specified together with gradients for three-dimensional beams).

**NF**

Number of different predefined field variables requested for any node (including field variables defined as initial conditions).

**TEMP (NT , N)**

An array containing the temperatures at the nodes involved in the constraint. This array is not used for a heat transfer, coupled temperature-displacement, or coupled thermal-electrical analysis since the temperatures are degrees of freedom of the problem.

**FIELD (NF , NT , N)**

An array containing all field variables at the nodes involved in the constraint.

**LTRAN (N)**

An integer array indicating whether the nodes in the MPC are transformed. If **LTRAN (I)=1**, a transformation is applied to node **I**; if **LTRAN (I)=0**, no transformation is applied.

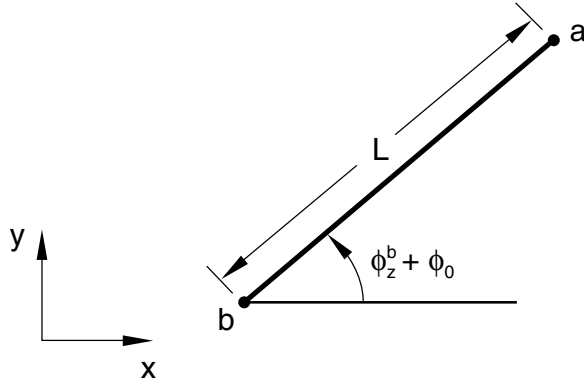
**TRAN (3 , 3 , N)**

An array containing the local-to-global transformation matrices for the nodes used in the MPC. If no transformation is present at node **I**, **TRAN (\* , \* , I)** is the identity matrix.

**Example: Nonlinear MPC**

As an example of a nonlinear MPC, consider the insertion of a rigid beam in a large-displacement, planar (two-dimensional) problem. This MPC is the two-dimensional version of library BEAM-type MPC. It can be implemented as a set of three different single degree of freedom MPCs or as a single nodal MPC. Here, the second method will be worked out because it is simpler and requires less data input.

Let  $a$  and  $b$  (see Figure 1.1.14–2) be the ends of the beam, with  $a$  the dependent end.



**Figure 1.1.14–2** Nonlinear MPC example: rigid beam.

The rigid beam will then define both components of displacement and the rotation at  $a$  in terms of the displacements and rotation at end  $b$  according to the set of equations:

$$\begin{aligned} f_1(\mathbf{u}^a, \mathbf{u}^b) &= x^a - x^b - L \cos(\phi_z^b + \phi_0) = 0, \\ f_2(\mathbf{u}^a, \mathbf{u}^b) &= y^a - y^b - L \sin(\phi_z^b + \phi_0) = 0, \\ f_3(\mathbf{u}^a, \mathbf{u}^b) &= \phi_z^a - \phi_z^b = 0, \end{aligned}$$

where  $\mathbf{u}^a = (u_x^a, u_y^a, \phi_z^a)$  and  $\mathbf{u}^b = (u_x^b, u_y^b, \phi_z^b)$ ,  $(x^a, y^a)$  and  $(x^b, y^b)$  are the current locations of  $a$  and  $b$ ,  $\phi_z^a$  and  $\phi_z^b$  are the rotations at  $a$  and  $b$  about the  $z$ -axis,  $L$  is the length of the link, and  $\phi_0$  is the original orientation of the link.

In terms of the original positions  $(X^a, Y^a)$  and  $(X^b, Y^b)$  of  $a$  and  $b$ ,

$$L = \sqrt{L_X^2 + L_Y^2}$$

and

$$\begin{aligned} \cos \phi_0 &= L_X / L, \\ \sin \phi_0 &= L_Y / L, \end{aligned}$$

where  $L_X = X^a - X^b$  and  $L_Y = Y^a - Y^b$ . Thus, the constraint equations can be expressed as

$$\begin{aligned}f_1(\mathbf{u}^a, \mathbf{u}^b) &= u_x^a - u_x^b + L_X - L_X \cos \phi_z^b + L_Y \sin \phi_z^b = 0, \\f_2(\mathbf{u}^a, \mathbf{u}^b) &= u_y^a - u_y^b + L_Y - L_X \sin \phi_z^b - L_Y \cos \phi_z^b = 0, \\f_3(\mathbf{u}^a, \mathbf{u}^b) &= \phi_z^a - \phi_z^b = 0.\end{aligned}$$

In light of the above formulation, the nontrivial portions of the matrices **A** and **JDOF** are

$$\mathbf{A}(1 : 3, 1 : 3, 1) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad \mathbf{A}(1 : 3, 1 : 3, 2) = \begin{pmatrix} -1 & 0 & L_X \sin \phi_z^b + L_Y \cos \phi_z^b \\ 0 & -1 & -L_X \cos \phi_z^b + L_Y \sin \phi_z^b \\ 0 & 0 & -1 \end{pmatrix}$$

and

$$\mathbf{JDOF}(1 : 3, 1) = \mathbf{JDOF}(1 : 3, 2) = \begin{pmatrix} 1 \\ 2 \\ 6 \end{pmatrix}.$$

Since degree of freedom 6 ( $\phi_z$ ) appears in this constraint, there must be an element in the mesh that uses that degree of freedom—a B21 beam element, for example.

If the above multi-point constraint is defined as type 1 with nodes  $a$  and  $b$ , the user subroutine **MPC** could be coded as follows:

```

SUBROUTINE MPC(UE,A,JDOF,MDOF,N,JTYPE,X,U,UINIT,MAXDOF,LMPC,
* KSTEP,KINC,TIME,NT,NF,TEMP,FIELD,LTRAN,TRAN)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION UE(MDOF),A(MDOF,MDOF,N),JDOF(MDOF,N),X(6,N),
* U(MAXDOF,N),UINIT(MAXDOF,N),TIME(2),TEMP(NT,N),
* FIELD(NF,NT,N),LTRAN(N),TRAN(3,3,N)
C
IF (JTYPE .EQ. 1) THEN
  COSFIB = COS(U(6,2))
  SINFIB = SIN(U(6,2))
  ALX = X(1,1) - X(1,2)
  ALY = X(2,1) - X(2,2)
C
  UE(1) = U(1,2) + ALX*(COSFIB-1.) - ALY*SINFIB
  UE(2) = U(2,2) + ALY*(COSFIB-1.) + ALX*SINFIB
  UE(3) = U(6,2)
C
  A(1,1,1) = 1.
  A(2,2,1) = 1.
  A(3,3,1) = 1.

```

```

A(1,1,2) = -1.
A(1,3,2) = ALX*SINFIB + ALY*COSFIB
A(2,2,2) = -1.
A(2,3,2) = -ALX*COSFIB + ALY*SINFIB
A(3,3,2) = -1.

C
JDOF(1,1) = 1
JDOF(2,1) = 2
JDOF(3,1) = 6
JDOF(1,2) = 1
JDOF(2,2) = 2
JDOF(3,2) = 6
END IF

C
RETURN
END

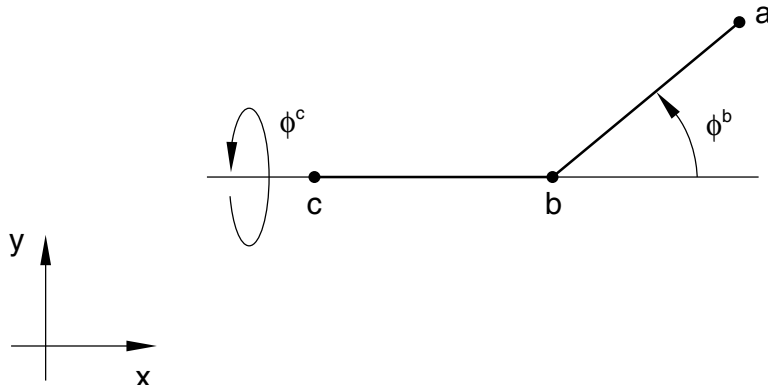
```

---

**Example: Nonlinear MPC involving finite rotations**


---

As an example of a nonlinear MPC involving finite rotations, consider a two-dimensional constant velocity joint that might be part of a robotics application. Let  $a$ ,  $b$ ,  $c$  (see Figure 1.1.14–3) be the nodes making up the joint, with  $a$  the dependent node.



**Figure 1.1.14–3** Nonlinear MPC example: constant velocity joint.

The joint is operated by prescribing an axial rotation  $\phi^c = \phi^c e_x$  at  $c$  and an out-of-plane rotation  $\phi^b = \phi^b e_z$  at  $b$ . The compounding of these two prescribed rotation fields will determine the total rotation at  $a$ . We can formally write this constraint as follows:

$$f(\phi^a, \phi^b, \phi^c) = \phi^a - \phi^b \circ \phi^c = 0,$$

where  $\circ$  denotes the rotation product. The formulation of the linearized constraint can be readily achieved from geometrically linear considerations in the deformed state.

In geometrically linear problems compound rotations are obtained simply as the linear superposition of individual rotation vectors. Consider the geometry depicted in Figure 1.1.14–3 and assume that the infinitesimal rotations  $\delta\theta^c = \delta\theta^c \mathbf{e}_x$  and  $\delta\theta^b = \delta\theta^b \mathbf{e}_z$  are applied at  $c$  and  $b$ , respectively. The rotation  $\delta\theta^a$  at  $a$  will simply be the sum of the vector  $\delta\theta^b$  to the vector  $\delta\theta^c$  rotated by an angle  $\phi^b$  about the  $z$ -axis. Thus, the linearized constraint can be written directly as

$$\begin{aligned}\delta f_1(\phi^a, \phi^b, \phi^c) &= \delta\theta_x^a - \cos(\phi^b)\delta\theta^c = 0, \\ \delta f_2(\phi^a, \phi^b, \phi^c) &= \delta\theta_y^a - \sin(\phi^b)\delta\theta^c = 0, \\ \delta f_3(\phi^a, \phi^b, \phi^c) &= \delta\theta_z^a - \delta\theta^b = 0.\end{aligned}$$

In light of this formulation, the nontrivial portions of the matrices **JDOF** and **A** are

$$\text{JDOF}(1 : 3, 1) = \begin{pmatrix} 4 \\ 5 \\ 6 \end{pmatrix}, \quad \text{JDOF}(1, 2) = 6, \quad \text{JDOF}(1, 3) = 4,$$

and

$$\mathbf{A}(1 : 3, 1 : 3, 1) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad \mathbf{A}(1 : 3, 1, 2) = \begin{pmatrix} 0 \\ 0 \\ -1 \end{pmatrix}, \quad \mathbf{A}(1 : 3, 1, 3) = \begin{pmatrix} -\cos(\phi^b) \\ -\sin(\phi^b) \\ 0 \end{pmatrix}.$$

Since degrees of freedom 4 ( $\phi_x$ ), 5 ( $\phi_y$ ), and 6 ( $\phi_z$ ) appear in this constraint, there must be an element in the mesh that uses these degrees of freedom—a B31 beam element, for example. The **MPC** subroutine has been coded with just this information. In this case Abaqus/Standard updates the dependent rotation field,  $\phi^a$ , based on the linearized constraint equations. Although the constraint is not satisfied exactly, good results are obtained as long as the rotation increments are kept small enough. A more rigorous derivation of the linearized constraint and the exact nonlinear recovery of the dependent degrees of freedom is presented in “Rotation variables,” Section 1.3.1 of the Abaqus Theory Manual.

If the above multi-point constraint is defined as type 1 with nodes  $a$ ,  $b$ , and  $c$ , user subroutine **MPC** could be coded as follows:

```

SUBROUTINE MPC (UE ,A ,JDOF ,MDOF ,N ,JTYPE ,X ,U ,UINIT ,MAXDOF ,LMPC ,
* KSTEP ,KINC ,TIME ,NT ,NF ,TEMP ,FIELD ,LTRAN ,TRAN)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION UE (MDOF) ,A (MDOF,MDOF,N) ,JDOF (MDOF,N) ,X (6,N) ,
* U (MAXDOF,N) ,UINIT (MAXDOF,N) ,TIME (2) ,TEMP (NT,N) ,

```

```
* FIELD (NF,NT,N) ,LTRAN (N) ,TRAN (3,3,N)
C
IF (JTYPE .EQ. 1) THEN
  A(1,1,1) = 1.
  A(2,2,1) = 1.
  A(3,3,1) = 1.
  A(3,1,2) = -1.
  A(1,1,3) = -COS(U(6,2))
  A(2,1,3) = -SIN(U(6,2))
C
  JDOF(1,1) = 4
  JDOF(2,1) = 5
  JDOF(3,1) = 6
  JDOF(1,2) = 6
  JDOF(1,3) = 4
END IF
C
RETURN
END
```

**1.1.15 ORIENT: User subroutine to provide an orientation for defining local material directions or local directions for kinematic coupling constraints or local rigid body directions for inertia relief.**

**Product:** Abaqus/Standard

## References

---

- “Orientations,” Section 2.2.5 of the Abaqus Analysis User’s Manual
- \*ORIENTATION
- “Eigenvalue analysis of a piezoelectric transducer,” Section 7.1.1 of the Abaqus Example Problems Manual

## Overview

---

User subroutine **ORIENT**:

- will be called at the start of the analysis at each location (material point, special-purpose element, coupling node, or reference point for inertia relief) for which local directions are defined with a user-subroutine-defined orientation;
- is used to define the direction cosines of a local system of (material) directions with respect to the default basis directions (default basis directions are defined as the global directions for continuum elements and as the default surface directions for shell, membrane, and surface elements, as described in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual);
- can be used to define the direction cosines orienting the layer of reinforcing material in membrane, shell, or surface elements (see “Defining reinforcement,” Section 2.2.3 of the Abaqus Analysis User’s Manual);
- can be used to provide a local system for defining the direction of action of rotary inertia, spring, dashpot, flexible joint, and elastic-plastic joint elements;
- can be used with gasket elements to define the local in-plane directions for three-dimensional area and three-dimensional link elements that consider transverse shear and membrane deformations (see “Defining the gasket behavior directly using a gasket behavior model,” Section 29.6.6 of the Abaqus Analysis User’s Manual);
- can be used to define a local system in which coupling constraints are applied (see “Coupling constraints,” Section 31.3.2 of the Abaqus Analysis User’s Manual, and “Kinematic coupling constraints,” Section 31.2.3 of the Abaqus Analysis User’s Manual);
- can be used to define a local system at the reference point for the rigid body directions in which inertia relief loads are applied for the entire model (see “Inertia relief,” Section 11.1.1 of the Abaqus Analysis User’s Manual);
- will ignore rotation angles defined for layers of composite solids (see “Solid (continuum) elements,” Section 25.1.1 of the Abaqus Analysis User’s Manual) but will take into account rotation angles

## ORIENT

defined for layers of composite shells (see “Using a shell section integrated during the analysis to define the section behavior,” Section 26.6.5 of the Abaqus Analysis User’s Manual, and “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual); and

- ignores any data specified for the associated orientation definition outside the user subroutine.

The local directions defined by user subroutine **ORIENT** must be specified relative to the default basis directions.

### User subroutine interface

---

```
SUBROUTINE ORIENT (T ,NOEL ,NPT ,LAYER ,KSPT ,COORDS ,BASIS ,
1 ORNAME ,NNODES ,CNODES ,JNNUM)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 ORNAME
C
DIMENSION T(3,3) ,COORDS(3) ,BASIS(3,3) ,CNODES(3,NNODES)
DIMENSION JNNUM(NNODES)

user coding to define T

RETURN
END
```

### Variable to be defined

---

#### T

An array containing the direction cosines of the preferred orientation in terms of the default basis directions. **T(1,1)**, **T(2,1)**, **T(3,1)** give the (1, 2, 3) components of the first direction; **T(1,2)**, **T(2,2)**, **T(3,2)** give the second direction; etc. For shell and membrane elements only the first and second directions are used. The directions do not have to be normalized. If the second direction is not orthogonal to the first direction, Abaqus/Standard will orthogonalize and normalize the second direction with respect to the first. The third direction is then determined by taking the cross product of the first and second directions. For planar elements the first two directions must lie in the plane of the element.

For use with coupling constraints (“Coupling constraints,” Section 31.3.2 of the Abaqus Analysis User’s Manual), the local basis directions are used as the local constraint directions for application of the kinematic constraint.

For use with inertia relief loads, the local basis directions are used as the rigid body direction vectors for computing the loads.

---

## Variables passed in for information

**NOEL**

Element number. This value is zero when the subroutine is called for use with coupling constraints or inertia relief loads.

**NPT**

Integration point number. This variable is set only for relevant uses.

**LAYER**

Layer number (for composite shells and layered solids). This variable is set only when relevant. It is equal to zero when it is irrelevant, such as in a regular solid element or in a shell element when transverse shear stiffness calculations are performed.

**KSPT**

Section point number within the current layer. This variable is set only when relevant. It is equal to zero when it is irrelevant, such as in a regular solid element or in a shell element when transverse shear stiffness calculations are performed.

**COORDS**

An array containing the initial coordinates of this point. This array contains the coordinates of the reference point for inertia relief loads.

**BASIS**

An array containing the direction cosines of the normal material basis directions in terms of the global coordinates in the original configuration. **BASIS(1,1)**, **BASIS(2,1)**, **BASIS(3,1)** give the 1-direction, etc. This is useful only in shells or membranes since in all other cases the basis is the global coordinate system.

**ORNAME**

User-specified orientation name, left justified, with one exception. When an overall section orientation is specified for a composite solid or shell section and the individual layer orientations are specified by an orientation angle, Abaqus defines an internal orientation name to represent the actual orientation of the layer. To avoid internal names, provide an orientation name rather than an orientation angle as part of the layer definition for each individual layer of a composite section.

**NNODES**

Number of element nodes. This value is two when the subroutine is called for use with a kinematic coupling definition, where the two nodes are the reference and current coupling node. When used with a distributing coupling definition, this number is equal to the number of coupling nodes plus one for the reference node. It is one when used with inertia relief loads since the local basis is defined at the reference point.

**CNODES**

An array containing the original coordinates of the nodes. When used with a kinematic coupling definition, the first entry defines the reference node coordinates, and the second entry defines the coupling node coordinates. When used with a distributing coupling definition, the first entry defines the reference node coordinates, and the subsequent entries define the coupling node coordinates in the order defined by the **JNNUM** array. When used with inertia relief loads, this array is not used. For all other uses the entry order follows that of the element definition node ordering.

**JNNUM**

An array containing the **NNODES** node numbers. When used with a kinematic coupling definition, the first entry is the reference node number, and the second entry is the node number for the current coupling node. When used with a distributing coupling definition, the first entry is the reference node number followed by the node numbers of all coupling nodes. When used with inertia relief loads, this array is not used. For all other uses the entry order follows that of the element definition node ordering.

## 1.1.16 RSURFU: User subroutine to define a rigid surface.

**Product:** Abaqus/Standard

### References

---

- “Analytical rigid surface definition,” Section 2.3.4 of the Abaqus Analysis User’s Manual
- \*SURFACE
- \*RIGID BODY
- “**RSURFU**,” Section 4.1.10 of the Abaqus Verification Manual

### Overview

---

User subroutine **RSURFU**:

- is used to define the surface of a rigid body for use in contact pairs;
- can be used to define a complex rigid surface if the various capabilities provided for defining a surface in Abaqus (see “Analytical rigid surface definition,” Section 2.3.4 of the Abaqus Analysis User’s Manual) are too restrictive;
- will be called at points on the slave surface of a contact pair or, if contact elements are used, at each integration point of each contact element with which the rigid surface is associated; and
- requires the definition of the closest point on the rigid surface, the normal and tangent directions, and the surface curvature.

### Overpenetration constraint

---

This routine must determine if a point on the slave surface has penetrated the rigid surface and define the local surface geometry. If the deforming and rigid surfaces are in contact at this point, Abaqus/Standard will impose a constraint at the point to prevent overpenetration. The local surface geometry must be defined to provide the necessary orientation for the constraint equations and friction directions and to allow Abaqus/Standard to compute the rate of change of these equations as the point moves around on the surface—the “tangent stiffness matrix” for the surface in the Newton algorithm. For the purpose of these calculations, it is best to define a smooth surface. If the surface is defined in a discontinuous manner, convergence may be adversely affected.

### Calculations to be performed

---

Each time **RSURFU** is called, Abaqus/Standard gives the current position of point  $A$  on the surface of the deforming structure,  $\mathbf{x}_A$ ; the current position of the rigid body reference point,  $\mathbf{x}_C$ ; the total displacements of both of these points,  $\mathbf{u}_A$  and  $\mathbf{u}_C$ ; and the total rotation of the rigid body reference point,  $\phi_C$ .

The routine should perform the following calculations:

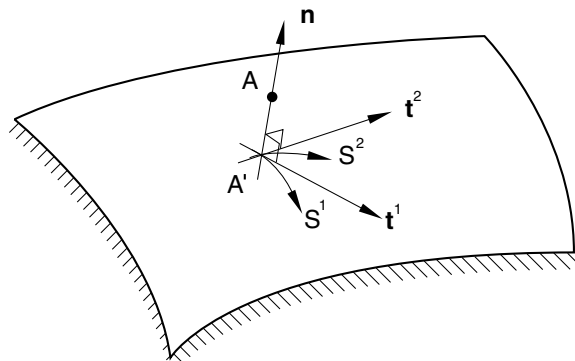
1. A point,  $A'$ , must be found on the rigid surface at which the normal to the surface passes through  $x_A$ . If there is not a unique point  $A'$ , the routine must choose the most suitable point (usually the closest  $A'$  to  $A$ ). The routine must pass back the coordinates of  $A'$  to Abaqus/Standard. For the surface-to-surface contact formulation, the slave normal, not the master normal, should be used.
2. **RSURFU** must define the distance,  $h$ , by which  $A$  has penetrated the surface below  $A'$ . A negative value of  $h$  means that  $A$  is outside the surface of the rigid body.
3. If the surfaces are in contact, which may sometimes be the case even if  $h$  is negative, **RSURFU** must define the local surface geometry.

### Defining the local surface geometry

---

There are two scenarios under which it is mandatory that the routine define the local surface geometry: if  $A$  has penetrated the surface— $h > 0$  if the surface behavior is truly rigid, or  $h$  is greater than the maximum overclosure value specified for modified surface behavior using either contact controls (see “Adjusting contact controls in Abaqus/Standard,” Section 32.3.6 of the Abaqus Analysis User’s Manual) or a modified pressure-overclosure relationship (see “Contact pressure-overclosure relationships,” Section 33.1.2 of the Abaqus Analysis User’s Manual)—and if  $A$  was in contact at the beginning of the increment, in which case the flag **LCLOSE**=1 (see the variable list for the definition of **LCLOSE**). The variable **LCLOSE** is not relevant for the surface-to-surface contact formulation and is always passed in as 0. The routine can be coded so that local surface geometry definitions are always provided regardless of the scenario.

The local surface geometry is specified by two orthogonal tangents to the rigid surface at  $A'$ , as well as the rates of change of the outward pointing normal at  $A'$ ,  $\mathbf{n}$ , with respect to local surface coordinates that are distance measuring along the tangents,  $S^1$  and  $S^2$  (see Figure 1.1.16–1).



**Figure 1.1.16–1** Local geometry on a rigid surface.

The tangents to the surface at  $A'$  must be defined so that their positive, right-handed cross product is the outward normal to the surface. For two-dimensional cases Abaqus/Standard assumes that the second tangent is  $(0, 0, -1)$ , so that when you give the direction cosines of the first tangent as  $(t_1, t_2, 0)$ , the

outward normal will be  $(-t_2, t_1, 0)$ . The rates of change of the normal with respect to  $S^1$  and  $S^2$  are required to define the local curvature of the surface.

## User subroutine interface

---

```

SUBROUTINE RSURFU(H,P,TGT,DNDS,X,TIME,U,CINAME,SLNAME,
1 MSNAME,NOEL,NODE,LCLOSE)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CINAME,SLNAME,MSNAME
C
DIMENSION P(3),TGT(3,2),DNDS(3,2),X(3,3),TIME(2),U(6,2)

```

*user coding to define H, P, TGT, and DNDS*

```

RETURN
END

```

## Variables to be defined

---

### H

Penetration of the point  $A$  on the deforming structure into the surface of the rigid body, measured down the outward normal to the rigid surface. A negative value of **H** indicates that  $A$  is outside the rigid surface. Even for a completely rigid surface,  $A$  may appear to penetrate the surface during the iterations because the kinematic constraints are not fully satisfied until an increment converges.

### P (3)

Position of the point  $A'$  on the surface of the rigid body closest to point  $A$  on the surface of the deforming structure.

### TGT (3, 2)

Direction cosines of the two unit tangents to the surface,  $t^1$  and  $t^2$ , at point  $A'$ . For two-dimensional cases only the first two components of  $t^1$  need be given since in this case Abaqus/Standard assumes that  $t^2$  is  $(0, 0, -1)$ .

### DNDS (3, 2)

Rates of change of the surface normal,  $\mathbf{n}$ , at  $A'$ , with respect to distance measuring coordinates,  $S^1$  and  $S^2$ , along  $t^1$  and  $t^2$ . For two-dimensional cases only the first two entries in the first column of **DNDS** ( $\partial n_1 / \partial S^1$ ,  $\partial n_2 / \partial S^1$ ) are required. The array **DNDS** is not required to be assigned for the surface-to-surface contact formulation.

---

**Variables passed in for information****X(K1,1)**

Current coordinates of point  $A$  on the surface of the deforming structure.

**X(K1,2)**

Current coordinates of the rigid body reference point.

**X(K1,3)**

Unit normal vector for point  $A$ ; relevant only for the surface-to-surface contact formulation.

**TIME(1)**

Value of step time at the end of the increment.

**TIME(2)**

Value of total time at the end of the increment.

**U(K1,1)**

Total displacement of point  $A$  on the surface of the deforming structure.

**U(K1,2)**

Total displacement and rotation of the rigid body reference point;  $k_1 = 1, 2, 3$  are the displacement components,  $k_1 = 4, 5, 6$  are the rotation components. For two-dimensional cases the only nonzero rotation component is  $k_1 = 6$ : **U(4,2)** and **U(5,2)** are both zero.

**CINAME**

User-specified surface interaction name, left justified. For user-defined contact elements it is either the element set name given for the interface definition or the optional name assigned to the interface definition.

**SLNAME**

Slave surface name. Passed in as blank if **RSURFU** is called for contact elements.

**MSNAME**

Master surface name. Passed in as blank if **RSURFU** is called for contact elements.

**NOEL**

Element label for contact elements. Passed in as zero if **RSURFU** is called for a contact pair.

**NODE**

Node number for point  $A$ . For the surface-to-surface contact formulation, this quantity is passed in as 0.

**LCLOSE**

Flag indicating contact status at the beginning of the increment. **LCLOSE**=1 indicates that  $A$  is in contact (closed) at the beginning of the increment. **LCLOSE**=0 indicates that  $A$  is not in contact (open)

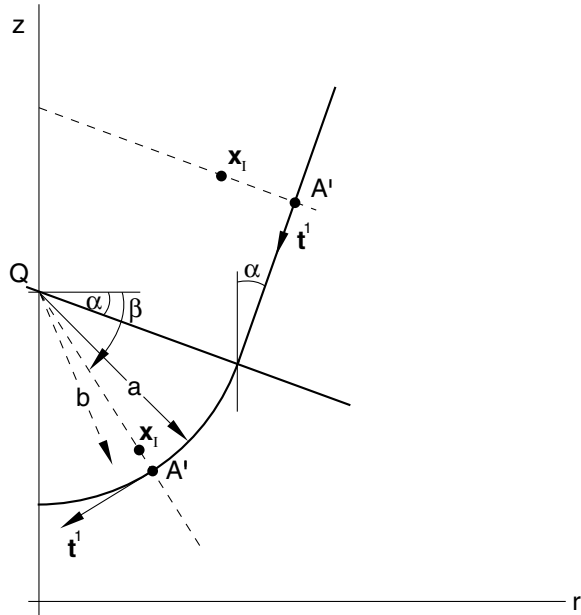
at the beginning of the increment. If **LCLOSE**=1, **P**, **TGT** and **DNDS** must be defined even if *A* opens during this increment. **LCLOSE** is not used for the surface-to-surface contact formulation and is passed in as 0.

### Example: Rigid punch

---

The input files for the following examples can be found in “**RSURFU**,” Section 4.1.10 of the Abaqus Verification Manual. The following discussion pertains only to the node-to-surface contact formulation.

Consider the punch shown in Figure 1.1.16–2.



**Figure 1.1.16–2** Cross-section of a rigid punch.

It consists of a spherical head of radius *a*, smoothly merging into a conical section with cone angle  $\alpha$ . The center of the sphere lies on the *z*-axis at *Q*. We assume that the punch is being driven down the *z*-axis by a prescribed displacement at the rigid body reference node defined as a boundary condition. (This same surface could be defined directly as a three-dimensional surface of revolution, as described in “Analytical rigid surface definition,” Section 2.3.4 of the Abaqus Analysis User’s Manual. We define it here in **RSURFU** as an illustration.)

A point (slave node) on the surface of the deforming body will be associated with the spherical head or with the conical part of the punch, depending on whether it lies above or below the cone that passes through *Q* and the circle of intersection of the sphere and cone. Thus, define

$$r = \sqrt{x_1^2 + x_2^2}, \quad z = x_3$$

in the three-dimensional case or

$$r = x_1, \quad z = x_2$$

in the axisymmetric case. Then, if  $r \tan \alpha < z_Q - z$ , the point is associated with the spherical surface. Otherwise, it is associated with the cone (both cases are indicated in Figure 1.1.16-2).

Consider first the axisymmetric case. Then, for  $r \tan \alpha < z_Q - z$  (the sphere) the overclosure is

$$h = a - b,$$

where

$$b = \sqrt{r^2 + (z - z_Q)^2}.$$

The position of the point  $A'$  on the rigid surface is  $(a \cos \beta, z_Q - a \sin \beta, 0)$ , where

$$\cos \beta = r/b, \quad \text{and} \quad \sin \beta = (z_Q - z)/b.$$

The tangent to the rigid surface at  $A'$  is  $\mathbf{t}^1 = (-\sin \beta, -\cos \beta, 0)$ . The positive direction for  $\mathbf{t}^1$  must be chosen so that the normal satisfies the right-hand rule with respect to  $\mathbf{t}^1$  and  $\mathbf{t}^2$  and points out of the rigid body. Also,  $dS^1 = a d\beta$ , so that

$$\frac{\partial \mathbf{n}}{\partial S^1} = \left( -\frac{1}{a} \sin \beta, -\frac{1}{a} \cos \beta, 0 \right).$$

For  $r \tan \alpha > z_Q - z$  (the conical surface) the clearance is

$$h = -r \cos \alpha + (z - z_Q) \sin \alpha + a,$$

and the position of the point  $A'$  on the rigid surface is  $(r + h \cos \alpha, z - h \sin \alpha, 0)$ . The surface tangent is  $\mathbf{t}^1 = (-\sin \alpha, -\cos \alpha, 0)$  and there is no change in  $\mathbf{n}$  with position, so that

$$\frac{\partial \mathbf{n}}{\partial S^1} = (0, 0, 0).$$

The routine can then be coded as follows:

```

SUBROUTINE RSURFU(H,P,TGT,DNDS,X,TIME,U,CINAME,SLNAME,
1      MSNAME,NOEL,NODE,LCLOSE)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CINAME,SLNAME,MSNAME
DIMENSION P(3),TGT(3,2),DNDS(3,2),X(3,2),TIME(2),U(6,2)
C
C      DEFINE THE FOLLOWING QUANTITIES:

```

```

C      A = RADIUS 'A' OF THE SPHERICAL HEAD
C      SINA = SINE (CONE ANGLE ALPHA)
C      COSA = COSINE (CONE ANGLE ALPHA)
C      Z0 = ORIGINAL 'Z' COORDINATE OF POINT 'Q'
C
C      A=5.0
C      SINA=0.5
C      COSA=0.86602
C      Z0=6.0
C      ZQ=Z0 + U(2,2)
C
C      TEST FOR SEGMENT
C
C      IF(X(1,1)*SINA/COSA.LT.ZQ-X(2,1))THEN
C
C      SPHERE
C
C      B=SQRT(X(1,1)**2 + (X(2,1)-ZQ)**2)
C      H=A-B
C      COSB=X(1,1)/B
C      SINB=(ZQ-X(2,1))/B
C      P(1)=A*COSB
C      P(2)=ZQ-A*SINB
C      TGT(1,1)=-SINB
C      TGT(2,1)=-COSB
C      DNDS(1,1)=-SINB/A
C      DNDS(2,1)=-COSB/A
C
C      ELSE
C
C      CONE
C
C      H=-X(1,1)*COSA+(X(2,1)-ZQ)*SINA+A
C      P(1)=X(1,1) + H*COSA
C      P(2)=X(2,1) - H*SINA
C      TGT(1,1)=-SINA
C      TGT(2,1)=-COSA
C      DNDS(1,1)=0.
C      DNDS(2,1)=0.
C
C      END IF
C
C      RETURN
C
C      END

```

The above case can be directly extended to three dimensions. For this purpose we assume that the radial axis,  $r$ , is in the global  $(x-y)$  plane, so that

$$r = \sqrt{x_1^2 + x_2^2}, \quad z = x_3.$$

For  $r \tan \alpha < z_Q - z$  (the sphere), the overclosure is  $h = a - b$ , where again

$$b = \sqrt{r^2 + (z - z_Q)^2}.$$

The point  $A'$  on the rigid surface is  $(a \cos \beta \cos \gamma, a \cos \beta \sin \gamma, z_Q - a \sin \beta)$ , where

$$\cos \gamma = \frac{x_1}{r}, \quad \sin \gamma = \frac{x_2}{r}.$$

For  $r = 0$ ,  $\gamma$  is not defined uniquely; in that case we arbitrarily choose  $\gamma = 0$ . We now need two tangents to the surface. The tangent  $\mathbf{t}^1$  used in the axisymmetric case is now

$$\mathbf{t}^1 = (-\sin \beta \cos \gamma, -\sin \beta \sin \gamma, -\cos \beta)$$

and the orthogonal tangent is

$$\mathbf{t}^2 = (-\sin \gamma, \cos \gamma, 0).$$

Again, the positive directions of  $\mathbf{t}^1$  and  $\mathbf{t}^2$  are chosen so that  $\mathbf{t}^1 \times \mathbf{t}^2$  defines an outward normal to the surface. The distance measures on the surface are

$$dS^1 = a d\beta, \quad dS^2 = a \cos \beta d\gamma$$

so that

$$\begin{aligned} \frac{\partial \mathbf{n}}{\partial S^1} &= \left( -\frac{1}{a} \sin \beta \cos \gamma, -\frac{1}{a} \sin \beta \sin \gamma, -\frac{1}{a} \cos \beta \right), \\ \frac{\partial \mathbf{n}}{\partial S^2} &= \left( -\frac{1}{a} \sin \gamma, \frac{1}{a} \cos \gamma, 0 \right). \end{aligned}$$

For the conical surface ( $r \tan \alpha \geq z_Q - z$ ), the surface separation is

$$h = -r \cos \alpha + (z - z_Q) \sin \alpha + a.$$

The point  $A'$  on the rigid surface is  $((r + h \cos \alpha) \cos \gamma, (r + h \cos \alpha) \sin \gamma, z - h \sin \alpha)$ ; and the surface tangents are

$$\begin{aligned} \mathbf{t}^1 &= (-\sin \alpha \cos \gamma, -\sin \alpha \sin \gamma, -\cos \alpha) \\ \mathbf{t}^2 &= (-\sin \gamma, \cos \gamma, 0). \end{aligned}$$

There is no change of  $\mathbf{n}$  with respect to  $S^1$ , and, in this case  $dS^2 = c d\gamma$ , where  $c = r + h \cos \alpha$  so that

$$\frac{\partial \mathbf{n}}{\partial S^2} = \left( -\frac{1}{c} \cos \alpha \sin \gamma, +\frac{1}{c} \cos \alpha \cos \gamma, 0 \right).$$

The routine can then be coded as follows:

```

      SUBROUTINE RSURFU(H,P,TGT,DNDS,X,TIME,U,CINAME,SLNAME,
1           MSNAME,NOEL,NODE,LCLOSE)
C
C       INCLUDE 'ABA_PARAM.INC'
C
C       CHARACTER*80 CINAME,SLNAME,MSNAME
C       DIMENSION P(3), TGT(3,2),DNDS(3,2), X(3,2), TIME(2), U(6,2)
C
C       DEFINE THE FOLLOWING QUANTITIES:
C       A = RADIUS 'A' OF THE SPHERICAL HEAD
C       SINA = SINE (CONE ANGLE ALPHA)
C       COSA = COSINE (CONE ANGLE ALPHA)
C       Z0 = ORIGINAL 'Z' COORDINATE OF POINT 'Q'
C
C       A=5.0
C       SINA=0.5
C       COSA=0.86603
C       Z0=5.0
C       ZQ= Z0 + U(3,2)
C
C       TEST FOR SEGMENT
C
C       R = SQRT(X(1,1)*X(1,1)+X(2,1)*X(2,1))
C       IF(R .GT. 0.0) THEN
C           COSG = X(1,1)/R
C           SING = X(2,1)/R
C       ELSE
C           COSG = 1.0
C           SING = 0.0
C       END IF
C       IF(R*SINA/COSA .LT. ZQ -X(3,1)) THEN
C
C           SPHERE
C
C           B=SQRT(R*R+(X(3,1)-ZQ)**2)
C           H=A-B
C           COSB=R/B
C           SINB=(ZQ-X(3,1))/B

```

```

P(1)=A*COSB*COSG
P(2)=A*COSB*SING
P(3)=ZQ-A*SINB
TGT(1,1)=-SINB*COSG
TGT(2,1)=-SINB*SING
TGT(3,1)=-COSB
TGT(1,2)=-SING
TGT(2,2)=COSG
TGT(3,2)=0.0
DNDS(1,1)=-SINB*COSG/A
DNDS(2,1)=-SINB*SING/A
DNDS(3,1)=-COSB/A
DNDS(1,2)=-SING/A
DNDS(2,2)=COSG/A
DNDS(3,2)=0.0
ELSE
C
C      CONE
C
H=-R*COSA+(X(3,1)-ZQ)*SINA+A
P(1)=(R+H*COSA)*COSG
P(2)=(R+H*COSA)*SING
P(3)=X(3,1)-H*SINA
TGT(1,1)=-SINA*COSG
TGT(2,1)=-SINA*SING
TGT(3,1)=-COSA
TGT(1,2)=-SING
TGT(2,2)=COSG
TGT(3,2)=0.0
DNDS(1,1)=0.0
DNDS(2,1)=0.0
DNDS(3,1)=0.0
C=R+H*COSA
DNDS(1,2)=-COSA*SING/C
DNDS(2,2)=COSA*COSG/C
DNDS(3,2)=0.0
END IF
C
RETURN
END

```

## 1.1.17 SDVINI: User subroutine to define initial solution-dependent state variable fields.

**Product:** Abaqus/Standard

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- \*INITIAL CONDITIONS
- “**SDVINI**,” Section 4.1.11 of the Abaqus Verification Manual

### Overview

---

User subroutine **SDVINI**:

- will be called for user-subroutine-defined initial solution-dependent state variable fields at particular material points, shell section points, contact slave nodes, or for user elements (see “Initial conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.2.1 of the Abaqus Analysis User’s Manual);
- can be used to initialize solution-dependent state variables allocated as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual; and
- returns a value of zero for any solution-dependent state variables that have no defined initial condition.

### Use of solution-dependent state variables in other user subroutines

---

Solution-dependent state variables initialized in **SDVINI** can be used and updated in the following user subroutines:

- **CREEP**
- **FRIC**
- **HETVAL**
- **UEL**
- **UEXPAN**
- **UGENS**
- **UHARD**
- **UMAT**
- **UMATHT**
- **USDFLD**
- **UTRS**

The solution-dependent state variables are passed into these routines in the order in which they are entered in **SDVINI**.

---

## User subroutine interface

---

```
SUBROUTINE SDVINI (STATEV, COORDS, NSTATV, NCRDS, NOEL, NPT,  
1 LAYER, KSPT)  
C  
INCLUDE 'ABA_PARAM.INC'  
C  
DIMENSION STATEV(NSTATV), COORDS(NCRDS)
```

*user coding to define STATEV (NSTATV)*

```
RETURN  
END
```

---

## Variables to be defined

---

**STATEV(1)**

First solution-dependent state variable.

**STATEV(2)**

Second solution-dependent state variable.

**STATEV(3)**

Third solution-dependent state variable.

**Etc.**

Only **NSTATV** solution-dependent state variable values should be defined.

---

## Variables passed in for information

---

**COORDS**

An array containing the initial coordinates of this point. Coordinates are not available for user elements.

**NSTATV**

User-defined number of solution-dependent state variables (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NCRDS**

Number of coordinates. This value is zero for user elements.

**NOEL**

Element number.

**NPT**

Integration point number in the element (not relevant for user elements).

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer or section. Section point 1 is used for all pure heat transfer and coupled temperature-displacement analyses.



## 1.1.18 SIGINI: User subroutine to define an initial stress field.

**Product:** Abaqus/Standard

### References

---

- “Initial conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.2.1 of the Abaqus Analysis User’s Manual
- \*INITIAL CONDITIONS

### Overview

---

User subroutine **SIGINI**:

- will be called for user-subroutine-defined initial stress fields at particular material points (these are the effective stress values for soils analysis);
- is called at the start of the analysis for each applicable material calculation point in the model; and
- can be used to define all active initial stress components at material points as functions of coordinates, element number, integration point number, etc.

### Stress components

---

The number of stress components that must be defined depends on the element type for which this call is being made. Part VI, “Elements,” of the Abaqus Analysis User’s Manual,” describes the element stresses. The order in which the components must be defined is the same as in the element definition. For example, in three-dimensional continuum elements six stress components must be defined in the order  $\sigma_{11}, \sigma_{22}, \sigma_{33}, \sigma_{12}, \sigma_{13}, \sigma_{23}$ .

### Initial stress field equilibrium

---

You should ensure that the initial stress field is in equilibrium with the applied forces and distributed loads by using a static step or a geostatic step to check the equilibrium of the initial stress field before starting the response history. See “Geostatic stress state,” Section 6.8.2 of the Abaqus Analysis User’s Manual, for a discussion of defining initial equilibrium conditions for problems that include pore fluid pressure.

### User subroutine interface

---

```

SUBROUTINE SIGINI(SIGMA,COORDS,NTENS,NCRDS,NOEL,NPT,LAYER,
1 KSPT,LREBAR,NAMES)
C
INCLUDE 'ABA_PARAM.INC'
C

```

```
DIMENSION SIGMA(NTENS),COORDS(NCRDS)
CHARACTER NAMES(2)*80
```

*user coding to define SIGMA(NTENS)*

```
RETURN
END
```

### **Variables to be defined**

---

#### **SIGMA(1)**

First stress component.

#### **SIGMA(2)**

Second stress component.

#### **SIGMA(3)**

Third stress component.

#### **Etc.**

Only **NTENS** stress values should be defined, where **NTENS** depends on the element type.

### **Variables passed in for information**

---

#### **COORDS**

An array containing the initial coordinates of this point.

#### **NTENS**

Number of stresses to be defined, which depends on the element type.

#### **NCRDS**

Number of coordinates.

#### **NOEL**

Element number.

#### **NPT**

Integration point number in the element.

#### **LAYER**

Layer number (for composite shells and layered solids).

#### **KSPT**

Section point number within the current layer.

**LREBAR**

Rebar flag. If **LREBAR**=1, the current integration point is associated with element rebar. Otherwise, **LREBAR**=0.

**NAMES (1)**

Name of the rebar to which the current integration point belongs, which is the name given in the rebar or rebar layer definition (“Defining reinforcement,” Section 2.2.3 of the Abaqus Analysis User’s Manual, or “Defining rebar as an element property,” Section 2.2.4 of the Abaqus Analysis User’s Manual). If no name was given in the rebar or rebar layer definition, this variable will be blank. This variable is relevant only when **LREBAR**=1.

**NAMES (2)**

Element type name (see Section EI.1, “Abaqus/Standard Element Index,” of the Abaqus Analysis User’s Manual).



## 1.1.19 UAMP: User subroutine to specify amplitudes.

**Product:** Abaqus/Standard

### References

---

- “Amplitude curves,” Section 30.1.2 of the Abaqus Analysis User’s Manual
- \*AMPLITUDE
- \*OUTPUT

### Overview

---

User subroutine **UAMP**:

- allows you to define the current value of an amplitude definition as a function of time;
- can be used to model control engineering aspects of your system when sensors are used (sensor values are from the beginning of the increment);
- can use a predefined number of state variables in their definition; and
- can optionally compute the derivatives and integrals of the amplitude function.

### Explicit solution dependence

---

The solution dependence introduced in this user subroutine is explicit: all data passed in the subroutine for information or to be updated are values at the beginning of that increment.

### User subroutine interface

---

```

SUBROUTINE UAMP(
*      ampName, time, ampValueOld, dt, nSvars, svars,
*      lFlagsInfo,
*      nSensor, sensorValues, sensorNames, jSensorLookUpTable,
*      AmpValueNew,
*      lFlagsDefine,
*      AmpDerivative, AmpSecDerivative, AmpIncIntegral,
*      AmpDoubleIntegral)
C
INCLUDE 'ABA_PARAM.INC'

C      time indices
parameter (iStepTime      = 1,
*              iTotalTime     = 2,
*              nTime          = 2)

```

```

C      flags passed in for information
      parameter (iInitialization    = 1,
*          iRegularInc      = 2,
*          iCuts            = 3
*          ikStep           = 4
*          nFlagsInfo       = 4)
C      optional flags to be defined
      parameter (iComputeDeriv     = 1,
*          iComputeSecDeriv   = 2,
*          iComputeInteg     = 3,
*          iComputeDoubleInteg = 4,
*          iStopAnalysis     = 5,
*          iConcludeStep     = 6,
*          nFlagsDefine      = 6)
      dimension time(nTime), lFlagsInfo(nFlagsInfo),
*          lFlagsDefine(nFlagsDefine)
      dimension jSensorLookUpTable(*)
      dimension sensorValues(nSensor), svarts(nSvarts)
      character*80 sensorNames(nSensor)
      character*80 ampName

user coding to define AmpValueNew, and
optionally lFlagsDefine, AmpDerivative, AmpSecDerivative,
AmpIncIntegral, AmpDoubleIntegral

      RETURN
      END

```

---

### Variable to be defined

#### **AmpValueNew**

Current value of the amplitude.

---

### Variables that can be updated

#### **lFlagsDefine**

Integer flag array to determine whether the computation of additional quantities is necessary or to set step continuation requirements.

#### **lFlagsDefine(iComputeDeriv)**

If set to 1, you must provide the computation of the amplitude derivative. The default is 0, which means that Abaqus computes the derivative automatically.

<b>lFlagsDefine(iComputeSecDeriv)</b>	
<b>lFlagsDefine(iComputeInteg)</b>	
<b>lFlagsDefine(iComputeDoubleInteg)</b>	
<b>lFlagsDefine(iStopAnalysis)</b>	
<b>lFlagsDefine(iConcludeStep)</b>	

If set to 1, you must provide the computation of the amplitude second derivative. The default is 0, which means that Abaqus computes the second derivative automatically.

If set to 1, you must provide the computation of the amplitude incremental integral. The default is 0, which means that Abaqus computes the incremental integral automatically.

If set to 1, you must provide the computation of the amplitude incremental double integral. The default is 0, which means that Abaqus computes the incremental integral automatically.

If set to 1, the analysis will be stopped and an error message will be issued. The default is 0, which means that Abaqus will not stop the analysis.

If set to 1, Abaqus will conclude the step execution and advance to the next step (if a next step exists). The default is 0.

### **svars**

An array containing the values of the solution-dependent state variables associated with this amplitude definition. The number of such variables is **nsvars** (see above). You define the meaning of these variables.

This array is passed into **UAMP** containing the values of these variables at the start of the current increment. In most cases they should be updated to be the values at the end of the increment.

### **AmpDerivative**

Current value of the amplitude derivative.

### **AmpSecDerivative**

Current value of the amplitude second derivative.

### **AmpIncIntegral**

Current value of the amplitude incremental integral.

### **AmpDoubleIntegral**

Current value of the amplitude incremental double integral.

## **Variables passed in for information**

---

### **ampName**

User-specified amplitude name, left justified.

**time (iStepTime)**

Current value of step time.

**time (iTTotalTime)**

Current value of total time.

**ampValueOld**

Old value of the amplitude from the previous increment.

**dt**

Current stable time increment.

**nSvars**

User-defined number of solution-dependent state variables associated with this amplitude definition.

**lFlagsInfo**

Integer flag array with information regarding the current call to **UAMP**.

**lFlagsInfo (iInitialization)**

This flag is equal to 1 if **UAMP** is called from the initialization phase of the first analysis step and is set to 0 otherwise.

**lFlagsInfo (iRegularInc)**

This flag is equal to 1 if **UAMP** is called from a regular increment and is set to 0 otherwise.

**lFlagsInfo (iCuts)**

Number of cutbacks in this increment.

**lFlagsInfo (ikStep)**

Step number.

**nSensor**

Total number of sensors in the model.

**sensorValues**

Array with sensor values at the end of the previous increment. Each sensor value corresponds to a history output variable associated with the output database request defining the sensor.

**sensorNames**

Array with user-defined sensor names in the entire model, left justified. Each sensor name corresponds to a sensor value provided with the output database request. All names will be converted to uppercase characters if lowercase or mixed-case characters were used in their definition.

**jSensorLookUpTable**

Variable that must be passed into the utility functions **IGETSENSORID** and **GETSENSORVALUE**.

---

**Example: Amplitude definition using sensor and state variables**

---

```

c      user amplitude subroutine
      Subroutine UAMP(
C          passed in for information and state variables
*          ampName, time, ampValueOld, dt, nSvars, svars,
*          lFlagsInfo,
*          nSensor, sensorValues, sensorNames, jSensorLookUpTable,
C          to be defined
*          ampValueNew,
*          lFlagsDefine,
*          AmpDerivative, AmpSecDerivative, AmpIncIntegral,
*          AmpDoubleIntegral)

      include 'aba_param.inc'

C      svars - additional state variables, similar to (V)UEL
      dimension sensorValues(nSensor), svars(nSvars)
      character*80 sensorNames(nSensor)
      character*80 ampName

C      time indices
      parameter( iStepTime      = 1,
*                  iTotTime       = 2,
*                  nTime          = 2)
C      flags passed in for information
      parameter( iInitialization = 1,
*                  iRegularInc   = 2,
*                  iCuts          = 3
*                  ikStep         = 4
*                  nFlagsInfo     = 4)
C      optional flags to be defined
      parameter( iComputeDeriv    = 1,
*                  iComputeSecDeriv = 2,
*                  iComputeInteg   = 3,
*                  iComputeDoubleInteg = 4,
*                  iStopAnalysis   = 5,
*                  iConcludeStep   = 6,
*                  nFlagsDefine    = 6)

      parameter( tStep=0.18d0, tAccelerateMotor = .00375d0,
*                  omegaFinal=23.26d0,

```

```

* zero=0.0d0, one=1.0d0, two=2.0d0, four=4.0d0)

dimension time(nTime), lFlagsInfo(nFlagsInfo),
*           lFlagsDefine(nFlagsDefine)
dimension jSensorLookUpTable(*)

lFlagsDefine(iComputeDeriv)      = 1
lFlagsDefine(iComputeSecDeriv)   = 1
lFlagsDefine(iComputeInteg)      = 1
lFlagsDefine(iComputeDoubleInteg) = 1

c      get sensor value
vTrans_CU1  = GetSensorValue('HORIZ_TRANSL_MOTION',
*                               jSensorLookUpTable,
*                               sensorValues)

if (ampName(1:22) .eq. 'MOTOR_WITH_STOP_SENSOR') then
  if (lFlagsInfo(iInitialization).eq.1) then
    AmpSecDerivative = zero
    AmpDerivative    = omegaFinal/tAccelerateMotor
    ampValueNew      = zero
    AmpIncIntegral   = zero
    AmpDoubleIntegral = zero

    svars(1) = zero
    svards(2) = zero
  else
    tim = time(iStepTime)

c      ramp up the angular rot velocity  of the
c      electric motor
c      after which hold constant
  if (tim .le. tAccelerateMotor) then
    AmpSecDerivative = zero
    AmpDerivative    = omegaFinal/tAccelerateMotor
    ampValueNew      = omegaFinal*tim/tAccelerateMotor
    AmpIncIntegral   = dt*(ampValueOld+ampValueNew) /
    two
    AmpDoubleIntegral = dt**2*(ampValueOld+ampValueNew) /
    four
  else
    AmpSecDerivative = zero

```

```

AmpDerivative      = zero
ampValueNew        = omegaFinal
AmpIncIntegral    = dt*(ampValueOld+ampValueNew) /
two
AmpDoubleIntegral = dt**2*(ampValueOld+ampValueNew) /
four
end if

c   retrieve old sensor value
vTrans_CU1_old = svars(1)

c   detect a zero crossing and count the number of crossings
if (vTrans_CU1_old*vTrans_CU1 .le. zero .and.
*      tim .gt. tAccelerateMotor ) then
    svars(2) = svars(2) + one
end if
nrCrossings = int(svars(2))

c   stop the motor if sensor crosses zero the second time
if (nrCrossings.eq.2) then
    ampValueNew = zero
    lFlagsDefine(iConcludeStep)=1
end if

c   store sensor value
svars(1) = vTrans_CU1

end if
end if

return
end

```



### 1.1.20 UANISOHYPER\_INV: User subroutine to define anisotropic hyperelastic material behavior using the invariant formulation.

**Product:** Abaqus/Standard

#### References

---

- “Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual
- \*ANISOTROPIC HYPERELASTIC
- “**UANISOHYPER\_INV** and **VUANISOHYPER\_INV**,” Section 4.1.13 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UANISOHYPER\_INV**:

- can be used to define the strain energy potential of anisotropic hyperelastic materials as a function of an irreducible set of scalar invariants;
- is called at all material calculation points of elements for which the material definition contains user-defined anisotropic hyperelastic behavior with an invariant-based formulation (“Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual);
- can include material behavior dependent on field variables or state variables; and
- requires that the values of the derivatives of the strain energy density function of the anisotropic hyperelastic material be defined with respect to the scalar invariants.

#### Enumeration of invariants

---

To facilitate coding and provide easy access to the array of invariants passed to user subroutine **UANISOHYPER\_INV**, an enumerated representation of each invariant is introduced. Any scalar invariant can, therefore, be represented uniquely by an *enumerated* invariant,  $I_n^*$ , where the subscript  $n$  denotes the order of the invariant according to the enumeration scheme in the following table:

Invariant	Enumeration, $n$
$\bar{I}_1$	1
$\bar{I}_2$	2
$J$	3
$\bar{I}_{4(\alpha\beta)}$	$4 + 2(\alpha - 1) + \beta(\beta - 1); \quad \alpha \leq \beta$
$\bar{I}_{5(\alpha\beta)}$	$5 + 2(\alpha - 1) + \beta(\beta - 1); \quad \alpha \leq \beta$

For example, in the case of three families of fibers there are a total of 15 invariants:  $\bar{I}_1$ ,  $\bar{I}_2$ ,  $J$ , six invariants of type  $\bar{I}_{4(\alpha\beta)}$ , and six invariants of type  $\bar{I}_{5(\alpha\beta)}$ , with  $\alpha, \beta = 1, 2, 3$  ( $\alpha \leq \beta$ ). The following correspondence exists between each of these invariants and their enumerated counterpart:

Enumerated invariant	Invariant
$I_1^*$	$\bar{I}_1$
$I_2^*$	$\bar{I}_2$
$I_3^*$	$J$
$I_4^*$	$\bar{I}_{4(11)}$
$I_5^*$	$\bar{I}_{5(11)}$
$I_6^*$	$\bar{I}_{4(12)}$
$I_7^*$	$\bar{I}_{5(12)}$
$I_8^*$	$\bar{I}_{4(22)}$
$I_9^*$	$\bar{I}_{5(22)}$
$I_{10}^*$	$\bar{I}_{4(13)}$
$I_{11}^*$	$\bar{I}_{5(13)}$
$I_{12}^*$	$\bar{I}_{4(23)}$
$I_{13}^*$	$\bar{I}_{5(23)}$
$I_{14}^*$	$\bar{I}_{4(33)}$
$I_{15}^*$	$\bar{I}_{5(33)}$

A similar scheme is used for the array **ZETA** of terms  $\zeta_{\alpha\beta} = \mathbf{A}_\alpha \cdot \mathbf{A}_\beta$ . Each term can be represented uniquely by an enumerated counterpart  $\zeta_m^*$ , as shown below:

Dot product	Enumeration, $m$
$\zeta_{\alpha\beta}$	$\alpha + \frac{1}{2}(\beta - 2)(\beta - 1); \quad \alpha < \beta$

As an example, for the case of three families of fibers there are three  $\zeta_{\alpha\beta}$  terms:  $\zeta_{12}$ ,  $\zeta_{13}$ , and  $\zeta_{23}$ . These are stored in the **ZETA** array as  $(\zeta_1^*, \zeta_2^*, \zeta_3^*)$ .

### Storage of arrays of derivatives of the energy function

---

The components of the array **UI1** of first derivatives of the strain energy potential with respect to the

scalar invariants,  $\partial U / \partial I_i^*$ , are stored using the enumeration scheme discussed above for the scalar invariants.

The elements of the array **UI2** of second derivatives of the strain energy function,  $\partial^2 U / \partial I_i^* \partial I_j^*$ , are laid out in memory using triangular storage: if  $k$  denotes the component in this array corresponding to the term  $\partial^2 U / \partial I_i^* \partial I_j^*$ , then  $k = i + j \times (j - 1)/2$ ; ( $i \leq j$ ). For example, the term  $\partial^2 U / \partial I_2^* \partial I_5^*$  is stored in component  $k = 2 + (5 \times 4)/2 = 12$  in the **UI2** array.

## **Special considerations for various element types**

---

There are several special considerations that need to be noted.

### **Shells that calculate transverse shear energy**

When **UANISOHYPER\_INV** is used to define the material response of shell elements that calculate transverse shear energy, Abaqus/Standard cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element's transverse shear stiffness. See "Shell section behavior," Section 26.6.4 of the Abaqus Analysis User's Manual, for guidelines on choosing this stiffness.

### **Elements with hourgassing modes**

When **UANISOHYPER\_INV** is used to define the material response of elements with hourgassing modes, you must define the hourglass stiffness for hourglass control based on the total stiffness approach. The hourglass stiffness is not required for enhanced hourglass control, but you can define a scaling factor for the stiffness associated with the drill degree of freedom (rotation about the surface normal). See "Section controls," Section 24.1.4 of the Abaqus Analysis User's Manual.

## **User subroutine interface**

---

```

SUBROUTINE UANISOHYPER_INV (AINV, UA, ZETA, NFIBERS, NINV,
1      UI1, UI2, UI3, TEMP, NOEL, CMNAME, INCMPFLAG, IHYBFLAG,
2      NUMSTATEV, STATEV, NUMFIELDV, FIELDV, FIELDVINC,
3      NUMPROPS, PROPS)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
DIMENSION AINV(NINV), UA(2),
2      ZETA(NFIBERS*(NFIBERS-1)/2)), UI1(NINV),
3      UI2(NINV*(NINV+1)/2), UI3(NINV*(NINV+1)/2),
4      STATEV(NUMSTATEV), FIELDV(NUMFIELDV),
5      FIELDVINC(NUMFIELDV), PROPS(NUMPROPS)

```

*user coding to define UA,UI1,UI2,UI3,STATEV*

```

RETURN
END

```

**Variables to be defined**

---

**UA(1)**

$U$ , strain energy density function. For a compressible material at least one derivative involving  $J$  should be nonzero. For an incompressible material all derivatives involving  $J$  are ignored.

**UA(2)**

$\tilde{U}_{dev}$ , the deviatoric part of the strain energy density of the primary material response. This quantity is needed only if the current material definition also includes Mullins effect (see “Mullins effect,” Section 19.6.1 of the Abaqus Analysis User’s Manual).

**UI1 (NINV)**

Array of derivatives of strain energy potential with respect to the scalar invariants,  $\partial U / \partial I_i^*$ , ordered using the enumeration scheme discussed above.

**UI2 (NINV\* (NINV+1) / 2)**

Array of second derivatives of strain energy potential with respect to the scalar invariants (using triangular storage),  $\partial^2 U / \partial I_i^* \partial I_j^*$ .

**UI3 (NINV\* (NINV+1) / 2)**

Array of derivatives with respect to  $J$  of the second derivatives of the strain energy potential (using triangular storage),  $\partial^3 U / \partial I_i^* \partial I_j^* \partial J$ . This quantity is needed only for compressible materials with a hybrid formulation (when **INCMPFLAG** = 0 and **IHYBFLAG** = 1).

**STATEV**

Array containing the user-defined solution-dependent state variables at this point. These are supplied as values at the start of the increment or as values updated by other user subroutines (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual) and must be returned as values at the end of the increment.

**Variables passed in for information**

---

**NFIBERS**

Number of families of fibers defined for this material.

**NINV**

Number of scalar invariants.

**TEMP**

Current temperature at this point.

**NOEL**

Element number.

**CMNAME**

User-specified material name, left justified.

**INCMPFLAG**

Incompressibility flag defined to be 1 if the material is specified as incompressible or 0 if the material is specified as compressible.

**IHYBFLAG**

Hybrid formulation flag defined to be 1 for hybrid elements; 0 otherwise.

**NUMSTATEV**

User-defined number of solution-dependent state variables associated with this material (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NUMFIELDV**

Number of field variables.

**FIELDV**

Array of interpolated values of predefined field variables at this material point at the end of the increment based on the values read in at the nodes (initial values at the beginning of the analysis and current values during the analysis).

**FIELDVINC**

Array of increments of predefined field variables at this material point for this increment, including any values updated by user subroutine **USDFLD**.

**NUMPROPS**

Number of material properties entered for this user-defined hyperelastic material.

**PROPS**

Array of material properties entered for this user-defined hyperelastic material.

**AINV (NINV)**

Array of scalar invariants,  $I_i^*$ , at each material point at the end of the increment. The invariants are ordered using the enumeration scheme discussed above.

**ZETA (NFIBERS\* (NFIBERS-1) / 2)**

Array of dot product between the directions of different families of fiber in the reference configuration,  $\zeta_{\alpha\beta} = \mathbf{A}_\alpha \cdot \mathbf{A}_\beta$ . The array contains the enumerated values  $\zeta_m^*$  using the scheme discussed above.

**Example: Anisotropic hyperelastic model of Kaliske and Schmidt**

As an example of the coding of user subroutine **UANISOHYPER\_INV**, consider the model proposed by Kaliske and Schmidt (2005) for nonlinear anisotropic elasticity with two families of fibers. The strain energy function is given by a polynomial series expansion in the form

$$U = \frac{1}{D}(J - 1)^2 + \sum_{i=1}^3 a_i(\bar{I}_1 - 3)^i + \sum_{j=1}^3 b_j(\bar{I}_2 - 3)^j + \sum_{k=2}^6 c_k(\bar{I}_{4(11)} - 1)^k + \sum_{l=2}^6 d_l(\bar{I}_{5(11)} - 1)^l \\ + \sum_{m=2}^6 e_m(\bar{I}_{4(22)} - 1)^m + \sum_{n=2}^6 f_n(\bar{I}_{5(22)} - 1)^n + \sum_{p=2}^6 g_p(\zeta_{12}\bar{I}_{4(12)} - \zeta_{12}^2)^p.$$

The code in user subroutine **UANISOHYPER\_INV** must return the derivatives of the strain energy function with respect to the scalar invariants, which are readily computed from the above expression. In this example auxiliary functions are used to facilitate enumeration of pseudo-invariants of type  $\bar{I}_{4(\alpha\beta)}$  and  $\bar{I}_{5(\alpha\beta)}$ , as well as for indexing into the array of second derivatives using symmetric storage. The user subroutine would be coded as follows:

```

subroutine uanisohyper_inv (aInv, ua, zeta, nFibers, nInv,
*                               ui1, ui2, ui3, temp, noel,
*                               cmname, incmpFlag, ihybFlag,
*                               numStatev, statev,
*                               numFieldv, fieldv, fieldvInc,
*                               numProps, props)
c
include 'aba_param.inc'
c
character *80 cmname
dimension aInv(nInv), ua(2), zeta(nFibers*(nFibers-1)/2)
dimension ui1(nInv), ui2(nInv*(nInv+1)/2)
dimension ui3(nInv*(nInv+1)/2), statev(numStatev)
dimension fieldv(numFieldv), fieldvInc(numFieldv)
dimension props(numProps)
c
parameter ( zero  = 0.d0,
*              one   = 1.d0,
*              two   = 2.d0,
*              three = 3.d0,
*              four  = 4.d0,
*              five  = 5.d0,
*              six   = 6.d0 )
c

```

```
C Kaliske-Schmidtt energy function (3D)
C
C      Read material properties
d=props(1)
dInv = one / d
a1=props(2)
a2=props(3)
a3=props(4)
b1=props(5)
b2=props(6)
b3=props(7)
c2=props(8)
c3=props(9)
c4=props(10)
c5=props(11)
c6=props(12)
d2=props(13)
d3=props(14)
d4=props(15)
d5=props(16)
d6=props(17)
e2=props(18)
e3=props(19)
e4=props(20)
e5=props(21)
e6=props(22)
f2=props(23)
f3=props(24)
f4=props(25)
f5=props(26)
f6=props(27)
g2=props(28)
g3=props(29)
g4=props(30)
g5=props(31)
g6=props(32)

C
C      Compute Udev and 1st and 2nd derivatives w.r.t invariants
C      - I1
bil = aInv(1)
term = bil-three
ua(2) = a1*term + a2*term**2 + a3*term***3
```

```

        ui1(1) = a1 + two*a2*term + three*a3*term**2
        ui2(indx(1,1)) = two*a2 + three*two*a3*term
C      - I2          bi2 = aInv(2)
        term = bi2-three
        ua(2) = ua(2) + b1*term + b2*term**2 + b3*term**3
        ui1(2) = b1 + two*b2*term + three*b3*term**2
        ui2(indx(2,2)) = two*b2 + three*two*b3*term
C      - I3 (=J)
        bi3 = aInv(3)
        term = bi3-one
        ui1(3) = two*dInv*term
        ui2(indx(3,3)) = two*dInv
C      - I4(11)
        nI411 = indxInv4(1,1)
        bi411 = aInv(nI411)
        term = bi411-one
        ua(2) = ua(2)
*           + c2*term**2 + c3*term**3 + c4*term**4
*           + c5*term**5 + c6*term**6
        ui1(nI411) =
*               two*c2*term
*               + three*c3*term**2
*               + four*c4*term**3
*               + five*c5*term**4
*               + six*c6*term**5
        ui2(indx(nI411,nI411)) =
*               two*c2
*               + three*two*c3*term
*               + four*three*c4*term**2
*               + five*four*c5*term**3
*               + six*five*c6*term**4
C      - I5(11)
        nI511 = indxInv5(1,1)
        bi511 = aInv(nI511)
        term = bi511-one
        ua(2) = ua(2)
*           + d2*term**2 + d3*term**3 + d4*term**4
*           + d5*term**5 + d6*term**6
        ui1(nI511) =
*               two*d2*term
*               + three*d3*term**2
*               + four*d4*term**3

```

```

*      + five*d5*term**4
*      + six*d6*term**5
ui2(indx(nI511,nI511)) =
*      two*d2
*      + three*two*d3*term
*      + four*three*d4*term**2
*      + five*four*d5*term**3
*      + six*five*d6*term**4

C - I4(22)
nI422 = indxInv4(2,2)
bi422 = aInv(nI422)
term = bi422-one
ua(2) = ua(2)
*      + e2*term**2 + e3*term**3 + e4*term**4
*      + e5*term**5 + e6*term**6
ui1(nI422) =
*      two*e2*term
*      + three*e3*term**2
*      + four*e4*term**3
*      + five*e5*term**4
*      + six*e6*term**5
ui2(indx(nI422,nI422)) =
*      two*e2
*      + three*two*e3*term
*      + four*three*e4*term**2
*      + five*four*e5*term**3
*      + six*five*e6*term**4

C - I5(22)
nI522 = indxInv5(2,2)
bi522 = aInv(nI522)
term = bi522-one
ua(2) = ua(2)
*      + f2*term**2 + f3*term**3 + f4*term**4
*      + f5*term**5 + f6*term**6
ui1(nI522) =
*      two*f2*term
*      + three*f3*term**2
*      + four*f4*term**3
*      + five*f5*term**4
*      + six*f6*term**5
ui2(indx(nI522,nI522)) =
*      two*f2

```

## UANISOHYPER\_INV

```

        *      + three*two*f3*term
        *      + four*three*f4*term**2
        *      + five*four*f5*term**3
        *      + six*five*f6*term**4
C     - I4(12)
nI412 = indxInv4(1,2)
bi412 = aInv(nI412)
term = zeta(1)*(bi412-zeta(1))
ua(2) = ua(2)
*      + g2*term**2 + g3*term**3
*      + g4*term**4 + g5*term**5
*      + g6*term**6
ui1(nI412) = zeta(1) * (
*      two*g2*term
*      + three*g3*term**2
*      + four*g4*term**3
*      + five*g5*term**4
*      + six*g6*term**5 )
ui2(indx(nI412,nI412)) = zeta(1)**2 * (
*      two*g2
*      + three*two*g3*term
*      + four*three*g4*term**2
*      + five*four*g5*term**3
*      + six*five*g6*term**4 )

C
c Add volumetric energy
c
        term = aInv(3) - one
        ua(1) = ua(2) + dInv*term*term
C
        return
        end
*
* Maps index from Square to Triangular storage of symmetric
        matrix integer function indx( i, j )
*
        include 'aba_param.inc'
*
        ii = min(i,j)
        jj = max(i,j)
*
        indx = ii + jj*(jj-1)/2

```

```
*  
    return  
end  
  
*  
*  
* Generate enumeration of Anisotropic Pseudo Invariants of  
type 4 integer function indxInv4( i, j )  
*  
    include 'aba_param.inc'  
  
*  
    ii = min(i,j)  
    jj = max(i,j)  
  
*  
    indxInv4 = 4 + jj*(jj-1) + 2*(ii-1)  
*  
    return  
end  
  
*  
*  
* Generate enumeration of Anisotropic Pseudo Invariants of  
type 5 integer function indxInv5( i, j )  
*  
    include 'aba_param.inc'  
  
*  
    ii = min(i,j)  
    jj = max(i,j)  
  
*  
    indxInv5 = 5 + jj*(jj-1) + 2*(ii-1)  
*  
    return  
end
```

---

## Additional reference

- Kaliske, M., and J. Schmidt, “Formulation of Finite Nonlinear Anisotropic Elasticity,” CADFEM GmbH Infoplaner 2/2005, vol. 2, pp. 22–23, 2005.



### 1.1.21 UANISOHYPER\_STRAIN: User subroutine to define anisotropic hyperelastic material behavior based on Green strain.

**Product:** Abaqus/Standard

#### References

---

- “Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual
- \*ANISOTROPIC HYPERELASTIC
- “UANISOHYPER\_INV and VUANISOHYPER\_INV,” Section 4.1.13 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UANISOHYPER\_STRAIN**:

- can be used to define the strain energy potential of anisotropic hyperelastic materials as a function of the components of the Green strain tensor;
- is called at all material calculation points of elements for which the material definition contains user-defined anisotropic hyperelastic behavior with a Green strain-based formulation (“Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual);
- can include material behavior dependent on field variables or state variables; and
- requires that the values of the derivatives of the strain energy density function of the anisotropic hyperelastic material be defined with respect to the components of the modified Green strain tensor.

#### Storage of strain components

---

In the array of modified Green strain, **E BAR**, direct components are stored first, followed by shear components. There are **NDI** direct and **NSHR** tensor shear components. The order of the components is defined in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual. Since the number of active stress and strain components varies between element types, the routine must be coded to provide for all element types with which it will be used.

#### Storage of arrays of derivatives of the energy function

---

The array of first derivatives of the strain energy function, **DU1**, contains **NTENS+1** components, with **NTENS=NDI+NSHR**. The first **NTENS** components correspond to the derivatives with respect to each component of the modified Green strain,  $\partial U / \partial \bar{\varepsilon}_{ij}^G$ . The last component contains the derivative with respect to the volume ratio,  $\partial U / \partial J$ .

The array of second derivatives of the strain energy function, **DU2**, contains **(NTENS+1) \* (NTENS+2) / 2** components. These components are ordered using the following triangular storage scheme:

<b>Component</b>	<b>2-D Case</b>	<b>3-D Case</b>
1	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{11}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{11}^G$
2	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{22}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{22}^G$
3	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{22}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{22}^G$
4	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{33}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{33}^G$
5	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{33}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{33}^G$
6	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{33}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{33}^G$
7	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{12}^G$
8	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{12}^G$
9	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{12}^G$
10	$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{12}^G$
11	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial J$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{13}^G$
12	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial J$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{13}^G$
13	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial J$	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{13}^G$
14	$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial J$	$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{13}^G$
15	$\partial^2 U / \partial J^2$	$\partial^2 U / \partial \bar{\varepsilon}_{13}^G \partial \bar{\varepsilon}_{13}^G$
16		$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{23}^G$
17		$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{23}^G$
18		$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{23}^G$
19		$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{23}^G$
20		$\partial^2 U / \partial \bar{\varepsilon}_{13}^G \partial \bar{\varepsilon}_{23}^G$
21		$\partial^2 U / \partial \bar{\varepsilon}_{23}^G \partial \bar{\varepsilon}_{23}^G$
22		$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial J$
23		$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial J$
24		$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial J$
25		$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial J$
26		$\partial^2 U / \partial \bar{\varepsilon}_{13}^G \partial J$

Component	2-D Case	3-D Case
27		$\partial^2 U / \partial \bar{\varepsilon}_{23}^G \partial J$
28		$\partial^2 U / \partial J^2$

Finally, the array of third derivatives of the strain energy function, **DU3**, also contains **(NTENS+1) \* (NTENS+2) / 2** components, each representing the derivative with respect to  $J$  of the corresponding component of **DU2**. It follows the same triangular storage scheme as **DU2**.

### Special considerations for various element types

---

There are several special considerations that need to be noted.

#### Shells that calculate transverse shear energy

When **UANISOHYPER\_STRAIN** is used to define the material response of shell elements that calculate transverse shear energy, Abaqus/Standard cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element's transverse shear stiffness. See "Shell section behavior," Section 26.6.4 of the Abaqus Analysis User's Manual, for guidelines on choosing this stiffness.

#### Elements with hourgassing modes

When **UANISOHYPER\_STRAIN** is used to define the material response of elements with hourgassing modes, you must define the hourglass stiffness for hourglass control based on the total stiffness approach. The hourglass stiffness is not required for enhanced hourglass control, but you can define a scaling factor for the stiffness associated with the drill degree of freedom (rotation about the surface normal). See "Section controls," Section 24.1.4 of the Abaqus Analysis User's Manual.

### User subroutine interface

---

```

SUBROUTINE UANISOHYPER_STRAIN (EBAR, AJ, UA, DU1, DU2, DU3,
1 TEMP, NOEL, CMNAME, INCMPLFLAG, IHYBFLAG, NDI, NSHR, NTENS,
2 NUMSTATEV, STATEV, NUMFIELDV, FIELDV, FIELDVINC,
3 NUMPROPS, PROPS)
C
      INCLUDE 'ABA_PARAM.INC'
C
      CHARACTER*80 CMNAME
C
      DIMENSION EBAR(NTENS), UA(2), DU1(NTENS+1),
2      DU2((NTENS+1)*(NTENS+2)/2),
3      DU3((NTENS+1)*(NTENS+2)/2),

```

## UANISOHYPER\_STRAIN

```
4      STATEV (NUMSTATEV) , FIELDV (NUMFIELDV) ,
5      FIELDVINC (NUMFIELDV) , PROPS (NUMPROPS)
```

*user coding to define UA,DU1,DU2,DU3,STATEV*

```
RETURN  
END
```

### Variables to be defined

---

#### UA (1)

$U$ , strain energy density function. For a compressible material at least one derivative involving  $J$  should be nonzero. For an incompressible material all derivatives involving  $J$  are ignored.

#### UA (2)

$\tilde{U}_{dev}$ , the deviatoric part of the strain energy density of the primary material response. This quantity is needed only if the current material definition also includes Mullins effect (see “Mullins effect,” Section 19.6.1 of the Abaqus Analysis User’s Manual).

#### DU1 (NTENS+1)

Derivatives of strain energy potential with respect to the components of the modified Green strain tensor,  $\partial U / \partial \bar{\varepsilon}_{ij}^G$ , and with respect to the volume ratio,  $\partial U / \partial J$ .

#### DU2 ( (NTENS+1) \* (NTENS+2) / 2 )

Second derivatives of strain energy potential with respect to the components of the modified Green strain tensor and the volume ratio (using triangular storage, as mentioned earlier).

#### DU3 ( (NTENS+1) \* (NTENS+2) / 2 )

Derivatives with respect to  $J$  of the second derivatives of the strain energy potential (using triangular storage, as mentioned earlier). This quantity is needed only for compressible materials with a hybrid formulation (when **INCPFLAG** = 0 and **IHYBFLAG** = 1).

#### STATEV

Array containing the user-defined solution-dependent state variables at this point. These are supplied as values at the start of the increment or as values updated by other user subroutines (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual) and must be returned as values at the end of the increment.

### Variables passed in for information

---

#### TEMP

Current temperature at this point.

**NOEL**

Element number.

**CMNAME**

User-specified material name, left justified.

**NDI**

Number of direct stress components at this point.

**NSHR**

Number of shear components at this point.

**NTENS**

Size of the stress or strain component array (**NDI** + **NSHR**).

**INCMPFLAG**

Incompressibility flag defined to be 1 if the material is specified as incompressible or 0 if the material is specified as compressible.

**IHYBFLAG**

Hybrid formulation flag defined to be 1 for hybrid elements; 0 otherwise.

**NUMSTATEV**

User-defined number of solution-dependent state variables associated with this material (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NUMFIELDV**

Number of field variables.

**FIELDV**

Array of interpolated values of predefined field variables at this material point at the end of the increment based on the values read in at the nodes (initial values at the beginning of the analysis and current values during the analysis).

**FIELDVINC**

Array of increments of predefined field variables at this material point for this increment, including any values updated by user subroutine **USDFLD**.

**NUMPROPS**

Number of material properties entered for this user-defined hyperelastic material.

**PROPS**

Array of material properties entered for this user-defined hyperelastic material.

## UANISOHYPER\_STRAIN

### EBAR (NTENS)

Modified Green strain tensor,  $\bar{\varepsilon}^G$ , at the material point at the end of the increment.

### AJ

$J$ , determinant of deformation gradient (volume ratio) at the end of the increment.

### Example: Orthotropic Saint-Venant Kirchhoff model

---

As a simple example of the coding of user subroutine **UANISOHYPER\_STRAIN**, consider the generalization to anisotropic hyperelasticity of the Saint-Venant Kirchhoff model. The strain energy function of the Saint-Venant Kirchhoff model can be expressed as a quadratic function of the Green strain tensor,  $\varepsilon^G$ , as

$$U(\varepsilon^G) = \frac{1}{2}\varepsilon^G : \mathbf{D} : \varepsilon^G,$$

where  $\mathbf{D}$  is the fourth-order elasticity tensor. The derivatives of the strain energy function with respect to the Green strain are given as

$$\frac{\partial U}{\partial \varepsilon^G} = \mathbf{D} : \varepsilon^G,$$

$$\frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G} = \mathbf{D}.$$

However, user subroutine **UANISOHYPER\_STRAIN** must return the derivatives of the strain energy function with respect to the modified Green strain tensor,  $\bar{\varepsilon}^G$ , and the volume ratio,  $J$ , which can be accomplished easily using the following relationship between  $\varepsilon^G$ ,  $\bar{\varepsilon}^G$ , and  $J$ :

$$\varepsilon^G = J^{\frac{2}{3}}\bar{\varepsilon}^G + \frac{1}{2}(J^{\frac{2}{3}} - 1)\mathbf{I},$$

where  $\mathbf{I}$  is the second-order identity tensor. Thus, using the chain rule we find

$$\frac{\partial U}{\partial \bar{\varepsilon}^G} = J^{\frac{2}{3}} \frac{\partial U}{\partial \varepsilon^G},$$

$$\frac{\partial U}{\partial J} = \frac{\partial \varepsilon^G}{\partial J} : \frac{\partial U}{\partial \varepsilon^G},$$

$$\frac{\partial^2 U}{\partial \bar{\varepsilon}^G \partial \bar{\varepsilon}^G} = J^{\frac{4}{3}} \frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G},$$

$$\frac{\partial^2 U}{\partial J^2} = \frac{\partial^2 \varepsilon^G}{\partial J^2} : \frac{\partial U}{\partial \varepsilon^G} + \frac{\partial \varepsilon^G}{\partial J} : \frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G} : \frac{\partial \varepsilon^G}{\partial J},$$

$$\frac{\partial^2 U}{\partial \bar{\varepsilon}^G \partial J} = \frac{2}{3J} J^{\frac{2}{3}} \frac{\partial U}{\partial \varepsilon^G} + J^{\frac{2}{3}} \frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G} : \frac{\partial \varepsilon^G}{\partial J},$$

where

$$\frac{\partial \varepsilon^G}{\partial J} = \frac{2}{3J} J^{\frac{2}{3}} (\bar{\varepsilon}^G + \frac{1}{2} \mathbf{I})$$

and

$$\frac{\partial^2 \varepsilon^G}{\partial J^2} = -\frac{1}{3J} \frac{\partial \varepsilon^G}{\partial J}.$$

In this example an auxiliary function is used to facilitate indexing into a fourth-order symmetric tensor. The user subroutine would be coded as follows:

```

subroutine uanisohyper_strain (
*      ebar, aj, ua, du1, du2, du3, temp, noel, cmname,
*      incmpFlag, ihybFlag, ndi, nshr, ntens,
*      numStatev, statev, numFieldv, fieldv, fieldvInc,
*      numProps, props)
c
      include 'aba_param.inc'
c
      dimension ebar(ntens), ua(2), du1(ntens+1)
      dimension du2((ntens+1)*(ntens+2)/2)
      dimension du3((ntens+1)*(ntens+2)/2)
      dimension statev(numStatev), fieldv(numFieldv)
      dimension fieldvInc(numFieldv), props(numProps)
c
      character*80 cmname
c
      parameter ( half      = 0.5d0,
$              one       = 1.d0,
$              two       = 2.d0,
$              third     = 1.d0/3.d0,
$              twothds   = 2.d0/3.d0,
$              four      = 4.d0 )
*
* Orthotropic Saint-Venant Kirchhoff strain energy function (3D)
*
      D1111=props(1)
      D1122=props(2)
      D2222=props(3)

```

## UANISOHYPER\_STRAIN

```
D1133=props(4)
D2233=props(5)
D3333=props(6)
D1212=props(7)
D1313=props(8)
D2323=props(9)
*
d2UdE11dE11 = D1111
d2UdE11dE22 = D1122
d2UdE11dE33 = D1133
*
d2UdE22dE11 = d2UdE11dE22
d2UdE22dE22 = D2222
d2UdE22dE33 = D2233
*
d2UdE33dE11 = d2UdE11dE33
d2UdE33dE22 = d2UdE22dE33
d2UdE33dE33 = D3333
*
d2UdE12dE12 = D1212
*
d2UdE13dE13 = D1313
*
d2UdE23dE23 = D2323
*
xpow = exp ( log(aj) * twothds )
detuInv = one / aj
*
E11 = xpow * ebar(1) + half * ( xpow - one )
E22 = xpow * ebar(2) + half * ( xpow - one )
E33 = xpow * ebar(3) + half * ( xpow - one )
E12 = xpow * ebar(4)
E13 = xpow * ebar(5)
E23 = xpow * ebar(6)
*
term1 = twothds * xpow * detuInv
dE11Dj = term1 * ( E11 + half )
dE22Dj = term1 * ( E22 + half )
dE33Dj = term1 * ( E33 + half )
dE12Dj = term1 * E12
dE13Dj = term1 * E13
dE23Dj = term1 * E23
```

```

term2 = - third * detuInv
d2E11DjDj = term2 * dE11Dj
d2E22DjDj = term2 * dE22Dj
d2E33DjDj = term2 * dE33Dj
d2E12DjDj = term2 * dE12Dj
d2E13DjDj = term2 * dE13Dj
d2E23DjDj = term2 * dE23Dj
*
dUdE11 = d2UdE11dE11 * E11
*      + d2UdE11dE22 * E22
*      + d2UdE11dE33 * E33
dUdE22 = d2UdE22dE11 * E11
*      + d2UdE22dE22 * E22
*      + d2UdE22dE33 * E33
dUdE33 = d2UdE33dE11 * E11
*      + d2UdE33dE22 * E22
*      + d2UdE33dE33 * E33
dUdE12 = two * d2UdE12dE12 * E12
dUdE13 = two * d2UdE13dE13 * E13
dUdE23 = two * d2UdE23dE23 * E23
*
U = half * ( E11*dUdE11 + E22*dUdE22 + E33*dUdE33 )
*      + E12*dUdE12 + E13*dUdE13 + E23*dUdE23
*
ua(2) = U
ua(1) = ua(2)
*
du1(1) = xpow * dUdE11
du1(2) = xpow * dUdE22
du1(3) = xpow * dUdE33
du1(4) = xpow * dUdE12
du1(5) = xpow * dUdE13
du1(6) = xpow * dUdE23
du1(7) = dUdE11*dE11Dj + dUdE22*dE22Dj + dUdE33*dE33Dj
*      + two * ( dUdE12*dE12Dj
*                  +dUdE13*dE13Dj
*                  +dUdE23*dE23Dj )
*
xpow2 = xpow * xpow
*
du2(indx(1,1)) = xpow2 * d2UdE11dE11
du2(indx(1,2)) = xpow2 * d2UdE11dE22

```

## UANISOHYPER\_STRAIN

```
du2(indx(2,2)) = xpow2 * d2UdE22dE22
du2(indx(1,3)) = xpow2 * d2UdE11dE33
du2(indx(2,3)) = xpow2 * d2UdE22dE33
du2(indx(3,3)) = xpow2 * d2UdE33dE33
du2(indx(1,4)) = zero
du2(indx(2,4)) = zero
du2(indx(3,4)) = zero
du2(indx(4,4)) = xpow2 * d2UdE12dE12
du2(indx(1,5)) = zero
du2(indx(2,5)) = zero
du2(indx(3,5)) = zero
du2(indx(4,5)) = zero
du2(indx(5,5)) = xpow2 * d2UdE13dE13
du2(indx(1,6)) = zero
du2(indx(2,6)) = zero
du2(indx(3,6)) = zero
du2(indx(4,6)) = zero
du2(indx(5,6)) = zero
du2(indx(6,6)) = xpow2 * d2UdE23dE23
*
du2(indx(1,7)) = term1 * dUdE11 + xpow * (
*      d2UdE11dE11 * dE11Dj
*      + d2UdE11dE22 * dE22Dj
*      + d2UdE11dE33 * dE33Dj )
du2(indx(2,7)) = term1 * dUdE22 + xpow * (
*      d2UdE22dE11 * dE11Dj
*      + d2UdE22dE22 * dE22Dj
*      + d2UdE22dE33 * dE33Dj )
du2(indx(3,7)) = term1 * dUdE33 + xpow * (
*      + d2UdE33dE11 * dE11Dj
*      + d2UdE33dE22 * dE22Dj
*      + d2UdE33dE33 * dE33Dj )
du2(indx(4,7)) = term1 * dUdE12
*      + xpow * two * d2UdE12dE12 * dE12Dj
du2(indx(5,7)) = term1 * dUdE13
*      + xpow * two * d2UdE13dE13 * dE23Dj
du2(indx(6,7)) = term1 * dUdE23
*      + xpow * two * d2UdE23dE23 * dE13Dj
du2(indx(7,7))= dUdE11*d2E11DjDj
*                  +dUdE22*d2E22DjDj
*                  +dUdE22*d2E22DjDj
*      + two*( dUdE12*d2E12DjDj
```

```
*           +dUdE13*d2E13DjDj
*           +dUdE23*d2E23DjDj)
*           + d2UdE11dE11 * dE11Dj * dE11Dj
*           + d2UdE22dE22 * dE22Dj * dE22Dj
*           + d2UdE33dE33 * dE33Dj * dE33Dj
*           + two   * ( d2UdE11dE22 * dE11Dj * dE22Dj
*                           +d2UdE11dE33 * dE11Dj * dE33Dj
*                           +d2UdE22dE33 * dE22Dj * dE33Dj )
*           + four  * ( d2UdE12dE12 * dE12Dj * dE12Dj
*           d2UdE13dE13 * dE13Dj * dE13Dj
*           d2UdE23dE23 * dE23Dj * dE23Dj )

*
*      return
*end
*
* Maps index from Square to Triangular storage
* of symmetric matrix
*
*      integer function indx( i, j )
*
*      include 'aba_param.inc'
*
*      ii = min(i,j)
*      jj = max(i,j)
*
*      indx = ii + jj*(jj-1)/2
*
*      return
*end
```



## 1.1.22 UCORR: User subroutine to define cross-correlation properties for random response loading.

**Product:** Abaqus/Standard

### References

---

- “Random response analysis,” Section 6.3.11 of the Abaqus Analysis User’s Manual
- \*CORRELATION
- “Random response to jet noise excitation,” Section 1.4.10 of the Abaqus Benchmarks Manual

### Overview

---

User subroutine **UCORR**:

- can be used to define the coefficients for the cross-correlation matrix in a random response analysis;
- will be called once for the combination of any two degrees of freedom with nonzero prescribed loads for each load case specified as a concentrated or distributed load or once for the combination of any two excitation directions specified as a base motion;
- allows correlation coefficients to be defined as a function of nodal coordinates; and
- ignores any data specified outside the user subroutine for the associated cross-correlation matrix.

### Cross-correlation for base motion excitation

---

The spatial correlation matrix for base motion excitation is defined by the coefficients  $\Psi_{ij}^{IJ}$  in user subroutine **UCORR**, where  $i, j$  are excitation directions and  $J$  corresponds to the  $J$ th frequency function referenced under load case  $I$ .

### Cross-correlation for point loads and distributed loads

---

The spatial correlation matrix of the load is defined as follows. Let  $F_{(N,i)}^I$  be the load applied to degree of freedom  $i$  at node  $N$  in load case  $I$ , through the use of a concentrated or distributed load. Let  $J$  correspond to the  $J$ th frequency function referenced under load case  $I$ . The spatial correlation matrix used in the random response analysis for this load case is then

$$\Psi_{(N,i)(M,j)}^{IJ} = C_{(N,i)(M,j)}^{IJ} F_{(N,i)}^I F_{(M,j)}^J,$$

where  $C_{(N,i)(M,j)}^{IJ}$  are the coefficients defined in user subroutine **UCORR**. Typically the load magnitude is given as 1.0; therefore, the load definition is simply selecting the nonzero terms that will appear in  $\Psi_{(N,i)(M,j)}^{IJ}$ .

**User subroutine interface**

---

```

SUBROUTINE UCORR(PSD,CORRR,CORRI,KSTEP,LCASE,JNODE1,JDOF1,
1 JNODE2,JDOF2,COOR1,COOR2)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION COOR1(3),COOR2(3)
CHARACTER*80 PSD

```

*user coding to define CORRR and CORRI*

```

RETURN
END

```

**Variables to be defined**

---

**CORRR**

Real part of the cross-correlation scaling factor.

**CORRI**

Imaginary part of the cross-correlation scaling factor.

**Variables passed in for information**

---

**PSD**

User-specified name for the frequency function that references this correlation, left justified.

**KSTEP**

Step number.

**LCASE**

Load case number,  $I$ .

**JNODE1**

First node involved,  $N$  (not used for base motion excitation).

**JDOF1**

Degree of freedom  $i$  at the first node (for concentrated or distributed load excitation) or global excitation direction  $i$  (for base motion excitation).

**JNODE2**

Second node involved,  $M$  (not used for base motion excitation).

**JDOF2**

Degree of freedom  $j$  at the second node (for concentrated or distributed load excitation) or global excitation direction  $j$  (for base motion excitation).

**COOR1**

An array containing the coordinates of the first node (not used for base motion excitation).

**COOR2**

An array containing the coordinates of the second node (not used for base motion excitation).



### 1.1.23 UDMGINI: User subroutine to define the damage initiation criterion.

**Product:** Abaqus/Standard

#### References

---

- “Progressive damage and failure,” Section 21.1.1 of the Abaqus Analysis User’s Manual
- “Modeling discontinuities as an enriched feature using the extended finite element method,” Section 10.6.1 of the Abaqus Analysis User’s Manual
- \*DAMAGE INITIATION

#### Overview

---

User subroutine **UDMGINI**:

- can be used to specify a user-defined damage initiation criterion;
- allows the specification of more than one failure mechanism in an element, with the most severe one governing the actual failure;
- can be used in combination with several Abaqus built-in damage evolution models, with each model corresponding to a particular failure mechanism;
- will be called at all integration points of elements for which the material definition contains user-defined damage initiation criterion;
- can call utility routine **GETVRM** to access material point data; and
- is currently available only for enriched elements.

#### User subroutine interface

---

```

SUBROUTINE UDMGINI (FINDEX,NFINDEX,FNORMAL,NDI,NSHR,NTENS,PROPS,
1 NPROPS,STATEV,INSTATEV,STRESS,STRAIN,STRAINEE,LXFEM,TIME,
2 DTIME,TEMP,DTEMP,PREDEF,DPRED,NFIELD,COORDS,NOEL,NPT,LAYER,
3 KSPT,KSTEP,KINC,KDIRCYC,KCYCLELCF,TIMECYC,SSE,SPD,SCD,SVD,
4 SMD,JMAC,JMATYP,MATLAYO,LACCFLA,CELENT,DROT,ORI)
C
INCLUDE 'ABA_PARAM.INC'

C
DIMENSION FINDEX(NFINDEX),FNORMAL(NDI,NFINDEX),COORDS(*),
1 STRESS(NTENS),STRAIN(NTENS),STRAINEE(NTENS),PROPS(NPROPS),
2 STATEV(INSTATV),PREDEF(NFIELD),DPRED(NFIELD),TIME(2),JMAC(*),
3 JMATYP(*),DROT(3,3),ORI(3,3)

```

*user coding to define **FINDEX**, and **FNORMAL***

```
RETURN
END
```

## Variables to be defined

---

### **FINDEX (NFINDEX)**

A Vector defining the indices for all the failure mechanisms.

### **FNORMAL (NDI , NFINDEX)**

An Array defining the normal direction to the fracture plane (three dimensions) or line (two dimensions) for each failure mechanism.

## Variables that can be updated

---

### **STATEV**

An array containing the user-defined solution-dependent state variables at this point. This array will be passed in containing the values of these variables at the start of the increment unless the values are updated in user subroutine **USDFLD**. They can be updated in this subroutine to their values at the end of the increment. You define the size of this array by allocating space for it (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information).

### **SSE , SPD , SCD , SVD , SMD**

Specific elastic strain energy, plastic dissipation, “creep” dissipation, viscous, and damage energy, respectively, passed in as the values at the start of the increment and should be updated to the corresponding specific energy values at the end of the increment. They have no effect on the solution, except that they are used for energy output.

## Variables passed in for information

---

### **NFINDEX**

Number of indices for all failure mechanisms.

### **NDI**

Number of direct stress components at this point.

### **NSHR**

Number of engineering shear stress components at this point.

### **NTENS**

Size of the stress or strain component array (**NRI** + **NSHR**).

### **PROPS (NPROPS)**

User-specified array of material constants associated with this user-defined failure criterion.

**NPROPS**

User-defined number of material constants associated with this user-defined failure criterion.

**NSTATV**

Number of solution-dependent state variables associated with this material (specified when space is allocated for the array; see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**STRESS (NTENS)**

An Array passed in as the current stress tensor. If a local orientation is used at the same point as user subroutine **UDMGINI**, the stress components will be in the local orientation; in the case of finite-strain analysis, the basis system in which stress components are stored rotates with the material.

**STRAIN (NTENS)**

An Array containing the current total strains. If a local orientation is used at the same point as user subroutine **UDMGINI**, the strain components will be in the local orientation; in the case of finite-strain analysis, the basis system in which strain components are stored rotates with the material.

**STRAINEE (NTENS)**

An Array containing the current elastic strains. If a local orientation is used at the same point as user subroutine **UDMGINI**, the elastic strain components will be in the local orientation; in the case of finite-strain analysis, the basis system in which elastic strain components are stored rotates with the material.

**LXFEM**

An integer flag to indicate an enriched element.

**TIME (1)**

Value of step time at the beginning of the current increment.

**TIME (2)**

Value of total time at the beginning of the current increment.

**DTIME**

Time increment.

**TEMP**

Temperature at the start of the increment.

**DTEMP**

Increment of temperature during the time increment.

**PREDEF**

An array containing the values of all of the user-specified predefined variables at this point at the start of the increment.

**DPRED**

An array containing the increments of all of the predefined variables during the time increment.

**NFIELD**

Number of user-specified predefined variables.

**COORDS**

An array containing the current coordinates of this point.

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**KSTEP**

Step number.

**KINC**

Increment number.

**KDIRCYC**

Iteration number in a direct cyclic analysis.

**KCYCLELCF**

Cycle number in a direct cyclic low-cycle fatigue analysis.

**TIMECYC**

Time period in one loading cycle in a direct cyclic analysis.

**JMAC**

Variable that must be passed into the **GETVRM** utility routine to access a material point variable.

**JMATYP**

Variable that must be passed into the **GETVRM** utility routine to access a material point variable.

**MATLATO**

Variable that must be passed into the **GETVRM** utility routine to access a material point variable.

**LACCFLA**

Variable that must be passed into the **GETVRM** utility routine to access a material point variable.

**CELENT**

Characteristic element length, which is a typical length of a line across an element for a first-order element; it is half of the same typical length for a second-order element. For beams and trusses it is a characteristic length along the element axis. For membranes and shells it is a characteristic length in the reference surface. For axisymmetric elements it is a characteristic length in the  $(r, z)$  plane only. For cohesive elements it is equal to the constitutive thickness.

**DROT (3, 3)**

Rotation increment matrix. This matrix represents the increment of rigid body rotation of the basis system in which the components of stress (STRESS) and strain (STRAIN) are stored. It is provided so that vector- or tensor-valued state variables can be rotated appropriately in this subroutine: stress and strain components are already rotated by this amount before **UDMGINI** is called. This matrix is passed in as a unit matrix for small-displacement analysis and for large-displacement analysis if the basis system for the material point rotates with the material (as in a shell element or when a local orientation is used).

**ORI (3, 3)**

Material orientation with respect to global basis.

**Example: User-defined damage initiation criterion with two different failure mechanisms**

---

As a simple example of the coding of user subroutine **UDMGINI**, consider a damage initiation criterion based on two different failure mechanisms: the maximum principal stress and the quadratic traction-interaction.

```

SUBROUTINE UDMGINI (FINDEX,NFINDEX,FNORMAL,NDI,NSHR,NTENS,PROPS,
 1 NPROPS,STATEV,NSTATEV,STRESS,STRAIN,STRAINEE,LXFEM,TIME,
 2 DTIME,TEMP,DTEMP,PREDEF,DPRED,NFIELD,COORDS,NOEL,NPT,
 3 KAYER,KSPT,KSTEP,INC,KDIRCYC,KCYCLELCF,TIMECYC,SSE,SPD,
 4 SCD,SVD,SMD,JMAC,JMATYP,MATLAYO,LACCFLA,CELENT,DROT,ORI)
C
INCLUDE 'ABA_PARAM.INC'
CC
DIMENSION FINDEX(NFINDEX),FNORMAL(NDI,NFINDEX),COORDS(*),
1 STRESS(NTENS),STRAIN(NTENS),STRAINEE(NTENS),PROPS(NPROPS),
2 STATEV(NSTATEV),PREDEF(NFIELD),DPRED(NFIELD),TIME(2),
3 JMAC(*),JMATYP(*),DROR(3,3),ORI(3,3)

DIMENSION PS(3), AN(3,3), WT(6)
PS(1)=0.0
PS(2)=0.0
PS(3)=0.0

```

```
C
C ROTATE THE STRESS TO GLOBAL SYSTEM IF THERE IS ORIENTATION
C
CALL ROTSIG(STRESS,ORI,WT,1,NDI,NSHR)
C
C MAXIMUM PRINCIPAL STRESS CRITERION
C
CALL SPRIND(WT,PS,AN,1,NDI,NSHR)
SIG1 = PS(1)
KMAX=1
DO K1 = 2, NDI
    IF(PS(K1).GT.SIG1) THEN
        SIG1 = PS(K1)
        KMAX = K1
    END IF
END DO
FINDEX(1) = SIG1/PROPS(1)
DO K1=1, NDI
    FNORMAL(K1,1) = AN(KMAX,K1)
END DO
C
C QUADRATIC TRACTION-INTERACTION CRITERION
C
FINDEX(2)=(STRESS(1)/PROPS(2))**2.0+(STRESS(NDI+1)/
$      PROPS(3))**2.0+(STRESS(NDI+2)/PROPS(4))**2.0
C
FINDEX(2)=sqrt(FINDEX(2))
C
DO K1=1, NDI
    FNORMAL(K1,2)=ORI(K1,1)
END DO
RETURN
END
```

### 1.1.24 UEL: User subroutine to define an element.

**Product:** Abaqus/Standard

*WARNING: This feature is intended for advanced users only. Its use in all but the simplest test examples will require considerable coding by the user/developer. “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual, should be read before proceeding.*

#### References

---

- “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual
- \*UEL PROPERTY
- \*USER ELEMENT

#### Overview

---

User subroutine **UEL**:

- will be called for each element that is of a general user-defined element type (i.e., not defined by a linear stiffness or mass matrix read either directly or from results file data) each time element calculations are required; and
- (or subroutines called by user subroutine **UEL**) must perform all of the calculations for the element, appropriate to the current activity in the analysis.

#### Wave kinematic data

---

For Abaqus/Aqua applications four utility routines—**GETWAVE**, **GETWAVEVEL**, **GETWINDVEL**, and **GETCURRVEL**—are provided to access the fluid kinematic data. These routines are used from within user subroutine **UEL** and are discussed in detail in “Obtaining wave kinematic data in an Abaqus/Aqua analysis,” Section 2.1.13.

#### User subroutine interface

---

```

SUBROUTINE UEL(RHS,AMATRX,SVARS,ENERGY,NDOFEL,NRHS,NSVARS,
1 PROPS,NPROPS,COORDS,MCRD,NNODE,U,DU,V,A,JTYPE,TIME,DTIME,
2 KSTEP,KINC,JELEM,PARAMS,NDLOAD,JDLTYP,ADLMAG,PREDEF,NPREDF,
3 LFLAGS,MLVARX,DDLMAG,MDLOAD,PNEWDT,JPROPS,NJPROP,PERIOD)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION RHS(MLVARX,*),AMATRX(NDOFEL,NDOFEL),PROPS(*),
1 SVARS(*),ENERGY(8),COORDS(MCRD,NNODE),U(NDOFEL),

```

```

2 DU (MLVARX,*) , V (NDOFEL) , A (NDOFEL) , TIME (2) , PARAMS (*),
3 JDLTYP (MDLOAD,*) , ADLMAG (MDLOAD,*) , DDLMAG (MDLOAD,*) ,
4 PREDEF (2 ,NPREFD ,NNODE) , LFLAGS (*) , JPROPS (*)

```

*user coding to define **RHS**, **AMATRX**, **SVARS**, **ENERGY**, and **PNEWDT***

```

RETURN
END

```

## Variables to be defined

---

These arrays depend on the value of the **LFLAGS** array.

### **RHS**

An array containing the contributions of this element to the right-hand-side vectors of the overall system of equations. For most nonlinear analysis procedures, **NRHS**=1 and **RHS** should contain the residual vector. The exception is the modified Riks static procedure (“Static stress analysis,” Section 6.2.2 of the Abaqus Analysis User’s Manual), for which **NRHS**=2 and the first column in **RHS** should contain the residual vector and the second column should contain the increments of external load on the element. **RHS (K1, K2)** is the entry for the **K1**th degree of freedom of the element in the **K2**th right-hand-side vector.

### **AMATRX**

An array containing the contribution of this element to the Jacobian (stiffness) or other matrix of the overall system of equations. The particular matrix required at any time depends on the entries in the **LFLAGS** array (see below).

All nonzero entries in **AMATRX** should be defined, even if the matrix is symmetric. If you do not specify that the matrix is unsymmetric when you define the user element, Abaqus/Standard will use the symmetric matrix defined by  $\frac{1}{2}([A] + [A]^T)$ , where  $[A]$  is the matrix defined as **AMATRX** in this subroutine. If you specify that the matrix is unsymmetric when you define the user element, Abaqus/Standard will use **AMATRX** directly.

### **SVARS**

An array containing the values of the solution-dependent state variables associated with this element. The number of such variables is **NSVARS** (see below). You define the meaning of these variables.

For general nonlinear steps this array is passed into **UEL** containing the values of these variables at the start of the current increment. They should be updated to be the values at the end of the increment, unless the procedure during which **UEL** is being called does not require such an update. This depends on the entries in the **LFLAGS** array (see below). For linear perturbation steps this array is passed into **UEL** containing the values of these variables in the base state. They should be returned containing perturbation values if you wish to output such quantities.

When **KINC** is equal to zero, the call to **UEL** is made for zero increment output (see “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). In this case the values returned will be used only for output purposes and are not updated permanently.

## **ENERGY**

For general nonlinear steps array **ENERGY** contains the values of the energy quantities associated with the element. The values in this array when **UEL** is called are the element energy quantities at the start of the current increment. They should be updated to the values at the end of the current increment. For linear perturbation steps the array is passed into **UEL** containing the energy in the base state. They should be returned containing perturbation values if you wish to output such quantities. The entries in the array are as follows:

<b>ENERGY (1)</b>	Kinetic energy.
<b>ENERGY (2)</b>	Elastic strain energy.
<b>ENERGY (3)</b>	Creep dissipation.
<b>ENERGY (4)</b>	Plastic dissipation.
<b>ENERGY (5)</b>	Viscous dissipation.
<b>ENERGY (6)</b>	“Artificial strain energy” associated with such effects as artificial stiffness introduced to control hourgassing or other singular modes in the element.
<b>ENERGY (7)</b>	Electrostatic energy.
<b>ENERGY (8)</b>	Incremental work done by loads applied within the user element.

When **KINC** is equal to zero, the call to **UEL** is made for zero increment output (see “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). In this case the energy values returned will be used only for output purposes and are not updated permanently.

---

## Variable that can be updated

### **PNEWDT**

Ratio of suggested new time increment to the time increment currently being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen). It is useful only during equilibrium iterations with the normal time incrementation, as indicated by **LFLAGS (3)=1**. During a severe discontinuity iteration (such as contact changes), **PNEWDT** is ignored unless CONVERT SDI=YES is specified for this step. The usage of **PNEWDT** is discussed below.

**PNEWDT** is set to a large value before each call to **UEL**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT × DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

## **Variables passed in for information**

---

### **Arrays:**

#### **PROPS**

A floating point array containing the **NPROPS** real property values defined for use with this element. **NPROPS** is the user-specified number of real property values. See “Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual.

#### **JPROPS**

An integer array containing the **NJPROP** integer property values defined for use with this element. **NJPROP** is the user-specified number of integer property values. See “Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual.

#### **COORDS**

An array containing the original coordinates of the nodes of the element. **COORDS (K1 , K2)** is the **K1**th coordinate of the **K2**th node of the element.

#### **U, DU, V, A**

Arrays containing the current estimates of the basic solution variables (displacements, rotations, temperatures, depending on the degree of freedom) at the nodes of the element at the end of the current increment. Values are provided as follows:

<b>U (K1)</b>	Total values of the variables. If this is a linear perturbation step, it is the value in the base state.
<b>DU (K1 , KRHS)</b>	Incremental values of the variables for the current increment for right-hand-side <b>KRHS</b> . If this is an eigenvalue extraction step, this is the eigenvector magnitude for eigenvector <b>KRHS</b> . For steady-state dynamics, <b>KRHS</b> = 1 denotes real components of perturbation displacement and <b>KRHS</b> = 2 denotes imaginary components of perturbation displacement.
<b>V (K1)</b>	Time rate of change of the variables (velocities, rates of rotation). Defined for implicit dynamics only ( <b>LFLAGS (1)</b> = 11 or 12).
<b>A (K1)</b>	Accelerations of the variables. Defined for implicit dynamics only ( <b>LFLAGS (1)</b> = 11 or 12).

**JDLTYP**

An array containing the integers used to define distributed load types for the element. Loads of type  $U_n$  are identified by the integer value  $n$  in **JDLTYP**; loads of type  $U_nNU$  are identified by the negative integer value  $-n$  in **JDLTYP**. **JDLTYP (K1 , K2)** is the identifier of the **K1**th distributed load in the **K2**th load case. For general nonlinear steps **K2** is always 1.

**ADLMAG**

For general nonlinear steps **ADLMAG (K1 , 1)** is the total load magnitude of the **K1**th distributed load at the end of the current increment for distributed loads of type  $U_n$ . For distributed loads of type  $U_nNU$ , the load magnitude is defined in **UEL**; therefore, the corresponding entries in **ADLMAG** are zero. For linear perturbation steps **ADLMAG (K1 , 1)** contains the total load magnitude of the **K1**th distributed load of type  $U_n$  applied in the base state. Base state loading of type  $U_nNU$  must be dealt with inside **UEL**. **ADLMAG (K1 , 2)**, **ADLMAG (K1 , 3)**, etc. are currently not used.

**DDLMAG**

For general nonlinear steps **DDLMAG** contains the increments in the magnitudes of the distributed loads that are currently active on this element for distributed loads of type  $U_n$ . **DDLMAG (K1 , 1)** is the increment of magnitude of the load for the current time increment. The increment of load magnitude is needed to compute the external work contribution. For distributed loads of type  $U_nNU$ , the load magnitude is defined in **UEL**; therefore, the corresponding entries in **DDLMAG** are zero. For linear perturbation steps **DDLMAG (K1 , K2)** contains the perturbation in the magnitudes of the distributed loads that are currently active on this element for distributed loads of type  $U_n$ . **K1** denotes the **K1**th perturbation load active on the element. **K2** is always 1, except for steady-state dynamics, where **K2=1** for real loads and **K2=2** for imaginary loads. Perturbation loads of type  $U_nNU$  must be dealt with inside **UEL**.

**PREDEF**

An array containing the values of predefined field variables, such as temperature in an uncoupled stress/displacement analysis, at the nodes of the element (“Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual).

The first index of the array, **K1**, is either 1 or 2, with 1 indicating the value of the field variable at the end of the increment and 2 indicating the increment in the field variable. The second index, **K2**, indicates the variable: the temperature corresponds to index 1, and the predefined field variables correspond to indices 2 and above. In cases where temperature is not defined, the predefined field variables begin with index 1. The third index, **K3**, indicates the local node number on the element.

<b>PREDEF (K1 , 1 , K3)</b>	Temperature.
<b>PREDEF (K1 , 2 , K3)</b>	First predefined field variable.
<b>PREDEF (K1 , 3 , K3)</b>	Second predefined field variable.
Etc.	Any other predefined field variable.

<b>PREDEF (K1 , K2 , K3)</b>	Total or incremental value of the <b>K2</b> th predefined field variable at the <b>K3</b> th node of the element.
<b>PREDEF (1 , K2 , K3)</b>	Values of the variables at the end of the current increment.
<b>PREDEF (2 , K2 , K3)</b>	Incremental values corresponding to the current time increment.

**PARAMS**

An array containing the parameters associated with the solution procedure. The entries in this array depend on the solution procedure currently being used when **UEL** is called, as indicated by the entries in the **LFLAGS** array (see below).

For implicit dynamics (**LFLAGS (1)** = 11 or 12) **PARAMS** contains the integration operator values, as:

<b>PARAMS (1)</b>	$\alpha$
<b>PARAMS (2)</b>	$\beta$
<b>PARAMS (3)</b>	$\gamma$

**LFLAGS**

An array containing the flags that define the current solution procedure and requirements for element calculations. Detailed requirements for the various Abaqus/Standard procedures are defined earlier in this section.

<b>LFLAGS (1)</b>	Defines the procedure type. See “Results file output format,” Section 5.1.2 of the Abaqus Analysis User’s Manual, for the key used for each procedure.
<b>LFLAGS (2)=0</b>	Small-displacement analysis.
<b>LFLAGS (2)=1</b>	Large-displacement analysis (nonlinear geometric effects included in the step; see “General and linear perturbation procedures,” Section 6.1.2 of the Abaqus Analysis User’s Manual).
<b>LFLAGS (3)=1</b>	Normal implicit time incrementation procedure. User subroutine <b>UEL</b> must define the residual vector in <b>RHS</b> and the Jacobian matrix in <b>AMATRX</b> .
<b>LFLAGS (3)=2</b>	Define the current stiffness matrix ( <b>AMATRX</b> = $K^{NM} = -\partial F^N / \partial u^M$ or $-\partial G^N / \partial u^M$ ) only.
<b>LFLAGS (3)=3</b>	Define the current damping matrix ( <b>AMATRX</b> = $C^{NM} = -\partial F^N / \partial \dot{u}^M$ or $-\partial G^N / \partial \dot{u}^M$ ) only.
<b>LFLAGS (3)=4</b>	Define the current mass matrix ( <b>AMATRX</b> = $M^{NM} = -\partial F^N / \partial \ddot{u}^M$ ) only. Abaqus/Standard always requests an initial mass matrix at the start of the analysis.

<b>LFLAGS (3)=5</b>	Define the current residual or load vector ( <b>RHS</b> = $F^N$ ) only.
<b>LFLAGS (3)=6</b>	Define the current mass matrix and the residual vector for the initial acceleration calculation (or the calculation of accelerations after impact).
<b>LFLAGS (3)=100</b>	Define perturbation quantities for output.
<b>LFLAGS (4)=0</b>	The step is a general step.
<b>LFLAGS (4)=1</b>	The step is a linear perturbation step.
<b>LFLAGS (5)=0</b>	The current approximations to $u^M$ , etc. were based on Newton corrections.
<b>LFLAGS (5)=1</b>	The current approximations were found by extrapolation from the previous increment.

**TIME (1)**

Current value of step time.

**TIME (2)**

Current value of total time.

**Scalar parameters:****DTIME**

Time increment.

**PERIOD**

Time period of the current step.

**NDOFEL**

Number of degrees of freedom in the element.

**MLVARX**

Dimensioning parameter used when several displacement or right-hand-side vectors are used.

**NRHS**

Number of load vectors. **NRHS** is 1 in most nonlinear problems: it is 2 for the modified Riks static procedure (“Static stress analysis,” Section 6.2.2 of the Abaqus Analysis User’s Manual), and it is greater than 1 in some linear analysis procedures and during substructure generation.

**NSVARS**

User-defined number of solution-dependent state variables associated with the element (“Defining the number of solution-dependent variables that must be stored within the element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**NPROPS**

User-defined number of real property values associated with the element (“Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**NJPROP**

User-defined number of integer property values associated with the element (“Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**MCRD**

**MCRD** is defined as the maximum of the user-defined maximum number of coordinates needed at any node point (“Defining the maximum number of coordinates needed at any nodal point” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual) and the value of the largest active degree of freedom of the user element that is less than or equal to 3. For example, if you specify that the maximum number of coordinates is 1 and the active degrees of freedom of the user element are 2, 3, and 6, **MCRD** will be 3. If you specify that the maximum number of coordinates is 2 and the active degrees of freedom of the user element are 11 and 12, **MCRD** will be 2.

**NNODE**

User-defined number of nodes on the element (“Defining the number of nodes associated with the element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**JTYPE**

Integer defining the element type. This is the user-defined integer value  $n$  in element type  $Un$  (“Assigning an element type key to a user-defined element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**KSTEP**

Current step number.

**KINC**

Current increment number.

**JELEM**

User-assigned element number.

**NDLOAD**

Identification number of the distributed load or flux currently active on this element.

**MDLOAD**

Total number of distributed loads and/or fluxes defined on this element.

**NPREFD**

Number of predefined field variables, including temperature. For user elements Abaqus/Standard uses one value for each field variable per node.

## UEL conventions

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The solution variables (displacement, velocity, etc.) are arranged on a node/degree of freedom basis. The degrees of freedom of the first node are first, followed by the degrees of freedom of the second node, etc.

## Usage with general nonlinear procedures

---

The values of  $u^N$  (and, in direct-integration dynamic steps,  $\dot{u}^N$  and  $\ddot{u}^N$ ) enter user subroutine **UEL** as their latest approximations at the end of the time increment; that is, at time  $t + \Delta t$ .

The values of  $H^\alpha$  enter the subroutine as their values at the beginning of the time increment; that is, at time  $t$ . It is your responsibility to define suitable time integration schemes to update  $H^\alpha$ . To ensure accurate, stable integration of internal state variables, you can control the time incrementation via **PNEWDT**.

The values of  $p^\beta$  enter the subroutine as the values of the total load magnitude for the  $\beta$ th distributed load at the end of the increment. Increments in the load magnitudes are also available.

In the following descriptions of the user element's requirements, it will be assumed that **LFLAGS (3)=1** unless otherwise stated.

### Static analysis (**LFLAGS (1)=1, 2**)

- $F^N = F^N(u^M, H^\alpha, p^\beta, t)$ .
- Automatic convergence checks are applied to the force residuals corresponding to degrees of freedom 1–7.
- You must define **AMATRX** =  $K^{NM} = -\partial F^N / \partial u^M$  and **RHS** =  $F^N$  and update the state variables,  $H^\alpha$ .

### Modified Riks static analysis (**LFLAGS (1)=1** and (**NRHS=2**)

- $F^N = F^N(u^M, H^\alpha, p^\beta)$ , where  $p^\beta = p_0^\beta + \lambda q^\beta$ ,  $p_0^\beta$  and  $q^\beta$  are fixed load parameters, and  $\lambda$  is the Riks (scalar) load parameter.
- Automatic convergence checks are applied to the force residuals corresponding to degrees of freedom 1–7.
- You must define **AMATRX** =  $K^{NM} = -\partial F^N / \partial u^M$ , **RHS (1)** =  $F^N$ , and **RHS (2)** =  $\Delta\lambda(\partial F^N / \partial \lambda)$  and update the state variables,  $H^\alpha$ . **RHS (2)** is the incremental load vector.

### Direct-integration dynamic analysis (**LFLAGS (1)=11, 12**)

- Automatic convergence checks are applied to the force residuals corresponding to degrees of freedom 1–7.
- **LFLAGS (3)=1**: Normal time increment. Either the Hilber-Hughes-Taylor or the backward Euler time integration scheme will be used. With  $\alpha$  set to zero for the backward Euler, both schemes imply

$$F^N = -M^{NM} \ddot{u}_{t+\Delta t} + (1 + \alpha) G^N_{t+\Delta t} - \alpha G_t^N,$$

where  $M^{NM} = M^{NM}(u^M, \dot{u}^M, H^\alpha, p^\beta, t, \dots)$  and  $G^N = G^N(u^M, \dot{u}^M, H^\alpha, p^\beta, t, \dots)$ ; that is, the highest time derivative of  $u^M$  in  $M^{NM}$  and  $G^N$  is  $\dot{u}^M$ , so that

$$-\frac{\partial F^N}{\partial \ddot{u}^M_{t+\Delta t}} = M^{NM}.$$

Therefore, you must store  $G_t^N$  as an internal state vector. If half-increment residual calculations are required, you must also store  $G_{t^-}^N$  as an internal state vector, where  $t^-$  indicates the time at the beginning of the previous increment. For  $\alpha = 0$ ,  $F^N = -M^{NM} \ddot{u}_{t+\Delta t} + G^N_{t+\Delta t}$  and  $G_t^N$  is not needed. You must define  $\text{AMATRX} = M^{NM}(\dot{u}/du) + (1 + \alpha) C^{NM}(\dot{u}/du) + (1 + \alpha) K^{NM}$ , where  $C^{NM} = -\partial G^N_{t+\Delta t}/\partial \dot{u}^M$  and  $K^{NM} = -\partial G^N_{t+\Delta t}/\partial u^M$ .  $\text{RHS} = F^N$  must also be defined and the state variables,  $H^\alpha$ , updated. Although the value of  $\alpha$  given in the dynamic step definition is passed into **UEL**, the value of  $\alpha$  can vary from element to element. For example,  $\alpha$  can be set to zero for some elements in the model where numerical dissipation is not desired.

- **LFLAGS (3)=5:** Half-increment residual ( $F_{1/2}^N$ ) calculation. Abaqus/Standard will adjust the time increment so that  $\max |F_{1/2}^N| < \text{tolerance}$  (where *tolerance* is specified in the dynamic step definition). The half-increment residual is defined as

$$F_{1/2}^N = -M^{NM} \ddot{u}_{t+\Delta t/2} + (1 + \alpha) G^N_{t+\Delta t/2} - \frac{\alpha}{2} (G_t^N + G_{t^-}^N),$$

where  $t^-$  indicates the time at the beginning of the previous increment ( $\alpha$  is a parameter of the Hilber-Hughes-Taylor time integration operator and will be set to zero if the backward Euler time integration operator is used). You must define  $\text{RHS} = F_{1/2}^N$ . To evaluate  $M^{NM}$  and  $G^N_{t+\Delta t/2}$ , you must calculate  $H^\alpha_{t+\Delta t/2}$ . These half-increment values will not be saved. **DTIME** will still contain  $\Delta t$  (not  $\Delta t/2$ ). The values contained in **U**, **V**, **A**, and **DU** are half-increment values.

- **LFLAGS (3)=4:** Velocity jump calculation. Abaqus/Standard solves  $-M^{NM} \Delta \dot{u}^M = 0$  for  $\Delta \dot{u}^M$ , so you must define  $\text{AMATRX} = M^{NM}$ .
- **LFLAGS (3)=6:** Initial acceleration calculation. Abaqus/Standard solves  $-M^{NM} \ddot{u}^M + G^N = 0$  for  $\ddot{u}^M$ , so you must define  $\text{AMATRX} = M^{NM}$  and  $\text{RHS} = G^N$ .

### Subspace-based dynamic analysis (**LFLAGS (1)=13**)

- The requirements are identical to those of static analysis, except that the Jacobian (stiffness), **AMATRX**, is not needed. No convergence checks are performed in this case.

### Quasi-static analysis (**LFLAGS (1)=21**)

- The requirements are identical to those of static analysis.

### Steady-state heat transfer analysis (**LFLAGS (1)=31**)

- The requirements are identical to those of static analysis, except that the automatic convergence checks are applied to the heat flux residuals corresponding to degrees of freedom 11, 12, ...

### Transient heat transfer analysis ( $\theta_{max}$ ) (**LFLAGS (1)=32 , 33**)

- Automatic convergence checks are applied to the heat flux residuals corresponding to degrees of freedom 11, 12, ...
- The backward difference scheme is always used for time integration; that is, Abaqus/Standard assumes that  $\dot{u}_{t+\Delta t} = \Delta u / \Delta t$ , where  $\Delta u = u_{t+\Delta t} - u_t$  and so  $d\dot{u}/du = 1/\Delta t$  always. For degrees of freedom 11, 12, ...,  $\max |\Delta u|$  will be compared against the user-prescribed maximum allowable nodal temperature change in an increment,  $\Delta\theta_{max}$ , for controlling the time integration accuracy.
- You need to define **AMATRX** =  $K^{NM} + (1/\Delta t) C^{NM}$ , where  $C^{NM}$  is the heat capacity matrix and **RHS** =  $F^N$ , and must update the state variables,  $H^\alpha$ .

### Geostatic analysis (**LFLAGS (1)=61**)

- Identical to static analysis, except that the automatic convergence checks are applied to the residuals corresponding to degrees of freedom 1–8.

### Steady-state coupled pore fluid diffusion/stress analysis (**LFLAGS (1)=62 , 63**)

- Identical to static analysis, except that the automatic convergence checks are applied to the residuals corresponding to degrees of freedom 1–8.

### Transient coupled pore fluid diffusion/stress (consolidation) analysis ( $u_w^{max}$ ) (**LFLAGS (1)=64 , 65**)

- Automatic convergence checks are applied to the residuals corresponding to degrees of freedom 1–8.
- The backward difference scheme is used for time integration; that is,  $\dot{u}_{t+\Delta t}^M = \Delta u^M / \Delta t$ , where  $\Delta u^M = u_{t+\Delta t}^M - u_t^M$ .
- For degree of freedom 8,  $\max |\Delta u^M|$  will be compared against the user-prescribed maximum wetting liquid pore pressure change,  $\Delta u_w^{max}$ , for automatic control of the time integration accuracy.
- You must define **AMATRX** =  $K^{NM} + (1/\Delta t) C^{NM}$ , where  $C^{NM}$  is the pore fluid capacity matrix and **RHS** =  $F^N$ , and must update the state variables,  $H^\alpha$ .

### Steady-state fully coupled thermal-stress analysis (**LFLAGS (1)=71**)

- Identical to static analysis, except that the automatic convergence checks are applied to the residuals corresponding to degrees of freedom 1–7 and 11, 12, ...

**Transient fully coupled thermal-stress analysis (  $\theta_{max}$  ) (**LFLAGS (1)=72 , 73**)**

- Automatic convergence checks are applied to the residuals corresponding to degrees of freedom 1–7 and 11, 12, ...
- The backward difference scheme is used for time integration; that is,  $\dot{u}_{t+\Delta t}^M = \Delta u^M / \Delta t$ , where  $\Delta u^M = u_{t+\Delta t}^M - u_t^M$ .
- For degrees of freedom 11, 12, ...,  $\max |\Delta u^M|$  will be compared against the user-prescribed maximum allowable nodal temperature change in an increment,  $\Delta\theta_{max}$ , for automatic control of the time integration accuracy.
- You must define **AMATRX** =  $K^{NM} + (1/\Delta t) C^{NM}$ , where  $C^{NM}$  is the heat capacity matrix and **RHS** =  $F^N$ , and must update the state variables,  $H^\alpha$ .

**Steady-state coupled thermal-electrical analysis (**LFLAGS (1)=75**)**

- The requirements are identical to those of static analysis, except that the automatic convergence checks are applied to the current density residuals corresponding to degree of freedom 9, in addition to the heat flux residuals.

**Transient coupled thermal-electrical analysis (  $\theta_{max}$  ) (**LFLAGS (1)=76 , 77**)**

- Automatic convergence checks are applied to the current density residuals corresponding to degree of freedom 9 and to the heat flux residuals corresponding to degree of freedom 11.
- The backward difference scheme is always used for time integration; that is, Abaqus/Standard assumes that  $\dot{u}_{t+\Delta t} = \Delta u / \Delta t$ , where  $\Delta u = u_{t+\Delta t} - u_t$ . Therefore,  $d\dot{u}/du = 1/\Delta t$  always. For degree of freedom 11  $\max |\Delta u|$  will be compared against the user-prescribed maximum allowable nodal temperature change in an increment,  $\Delta\theta_{max}$ , for controlling the time integration accuracy.
- You must define **AMATRX** =  $K^{NM} + (1/\Delta t) C^{NM}$ , where  $C^{NM}$  is the heat capacity matrix and **RHS** =  $F^N$ , and must update the state variables,  $H^\alpha$ .

---

**Usage with linear perturbation procedures**

“General and linear perturbation procedures,” Section 6.1.2 of the Abaqus Analysis User’s Manual, describes the linear perturbation capabilities in Abaqus/Standard. Here, base state values of variables will be denoted by  $u^M$ ,  $H^\alpha$ , etc. Perturbation values will be denoted by  $\tilde{u}^M$ ,  $\tilde{H}^\alpha$ , etc.

Abaqus/Standard will not call user subroutine **UEL** for the eigenvalue buckling prediction procedure.

For response spectrum, random response, transient modal dynamic, and mode-based steady-state dynamic procedures, user subroutine **UEL** is called only in a prior natural frequency extraction analysis, and the mass and stiffness contributions are taken into account during modal superposition.

For direct-solution and mode-based steady-state dynamic, complex eigenvalue extraction, matrix generation, and substructure generation procedures, Abaqus/Standard will call user subroutine **UEL**, but only mass and stiffness contributions will be taken into account. The damping contributions will be neglected.

**Static analysis (**LFLAGS (1)=1, 2**)**

- Abaqus/Standard will solve  $K^{NM} \tilde{u}^M = \tilde{P}^N$  for  $\tilde{u}^M$ , where  $K^{NM}$  is the base state stiffness matrix and the perturbation load vector,  $\tilde{P}^N$ , is a linear function of the perturbation loads,  $\tilde{p}$ ; that is,  $\tilde{P}^N = (\partial F / \partial \tilde{p}) \tilde{p}$ .
- **LFLAGS (3)=1:** You must define **AMATRX** =  $K^{NM}$  and **RHS** =  $\tilde{P}^N$ .
- **LFLAGS (3)=100:** You must compute perturbations of the internal variables,  $\tilde{H}^\alpha$ , and define **RHS** =  $\tilde{P}^N - K^{NM} \tilde{u}^M$  for output purposes.

**Eigenfrequency extraction analysis (**LFLAGS (1)=41**)**

- $F^N = -M^{NM} \ddot{\tilde{u}} + G^N(u^M + \tilde{u}^M, \dots) = -M^{NM} \ddot{\tilde{u}} + (\partial G^N / \partial u^M) \tilde{u}^M$ .
- Abaqus/Standard will solve  $K^{NM} \phi_i^M = \omega_i^2 M^{NM} \phi_i^M$  for  $\phi_i^N$  and  $\omega_i$ , where  $K^{NM} = -\partial F^N / \partial u^M$  is the base state stiffness matrix and  $M^{NM} = -\partial F^{NM} / \partial \ddot{u}^M$  is the base state mass matrix.
- **LFLAGS (3)=2:** Define **AMATRX** =  $K^{NM}$ .
- **LFLAGS (3)=4:** Define **AMATRX** =  $M^{NM}$ .

**Example: Structural and heat transfer user element**

---

Both a structural and a heat transfer user element have been created to demonstrate the usage of subroutine **UEL**. These user-defined elements are applied in a number of analyses. The following excerpt is from the verification problem that invokes the structural user element in an implicit dynamics procedure:

```
*USER ELEMENT, NODES=2, TYPE=U1, PROPERTIES=4, COORDINATES=3,
VARIABLES=12
1, 2, 3
*ELEMENT, TYPE=U1
101, 101, 102
*ELGEN, ELSET=UTRUSS
101, 5
*UEL PROPERTY, ELSET=UTRUSS
0.002, 2.1E11, 0.3, 7200.
```

The user element consists of two nodes that are assumed to lie parallel to the  $x$ -axis. The element behaves like a linear truss element. The supplied element properties are the cross-sectional area, Young's modulus, Poisson's ratio, and density, respectively.

The next excerpt shows the listing of the subroutine. The user subroutine has been coded for use in a perturbation static analysis; general static analysis, including Riks analysis with load incrementation defined by the subroutine; eigenfrequency extraction analysis; and direct-integration dynamic analysis. The names of the verification input files associated with the subroutine and these procedures can be found in “**UEL**,” Section 4.1.14 of the Abaqus Verification Manual. The subroutine performs all calculations required for the relevant procedures as described earlier in this section. The flags passed in through the **LFLAGS** array are used to associate particular calculations with solution procedures.

During a modified Riks analysis all force loads must be passed into **UEL** by means of distributed load definitions such that they are available for the definition of incremental load vectors; the load keys **Un** and **UnNU** must be used properly, as discussed in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual. The coding in subroutine **UEL** must distribute the loads into consistent equivalent nodal forces and account for them in the calculation of the **RHS** and **ENERGY** arrays.

```

SUBROUTINE UEL (RHS ,AMATRX ,SVARS ,ENERGY ,NDOFEL ,NRHS ,NSVARS ,
1      PROPS ,NPROPS ,COORDS ,MCRD ,NNODE ,U ,DU ,V ,A ,JTYPE ,TIME ,
2      DTIME ,KSTEP ,KINC ,JELEM ,PARAMS ,NDLOAD ,JDLTYP ,ADLMAG ,
3      PREDEF ,NPREFD ,LFLAGS ,MLVARX ,DDLMAG ,MDLOAD ,PNEWDT ,
4      JPROPS ,NJPROP ,PERIOD)

C
INCLUDE 'ABA_PARAM.INC'
PARAMETER ( ZERO = 0.D0 , HALF = 0.5D0 , ONE = 1.D0 )

C
DIMENSION RHS (MLVARX,*),AMATRX (NDOFEL,NDOFEL),
1      SVARS (NSVARS) ,ENERGY (8) ,PROPS (*) ,COORDS (MCRD ,NNODE) ,
2      U (NDOFEL) ,DU (MLVARX,*),V (NDOFEL) ,A (NDOFEL) ,TIME (2) ,
3      PARAMS (3) ,JDLTYP (MDLOAD ,*) ,ADLMAG (MDLOAD ,*) ,
4      DDLMAG (MDLOAD ,*) ,PREDEF (2,NPREFD,NNODE) ,LFLAGS (*) ,
5      JPROPS (*)
DIMENSION SRESID (6)

C
C UEL SUBROUTINE FOR A HORIZONTAL TRUSS ELEMENT
C
C      SRESID - stores the static residual at time t+dt
C      SVARS - In 1-6, contains the static residual at time t
C                  upon entering the routine. SRESID is copied to
C                  SVARS(1-6) after the dynamic residual has been
C                  calculated.
C      - For half-increment residual calculations: In 7-12,
C          contains the static residual at the beginning
C          of the previous increment. SVARS(1-6) are copied
C          into SVARS(7-12) after the dynamic residual has
C          been calculated.

C
AREA = PROPS (1)
E    = PROPS (2)
ANU = PROPS (3)
RHO = PROPS (4)

C
ALEN = ABS (COORDS (1,2)-COORDS (1,1))
AK   = AREA*E/ALEN

```

```

AM      = HALF*AREA*RHO*ALEN
C
DO K1 = 1, NDOFEL
  SRESID(K1) = ZERO
  DO KRHS = 1, NRHS
    RHS(K1,KRHS) = ZERO
  END DO
  DO K2 = 1, NDOFEL
    AMATRX(K2,K1) = ZERO
  END DO
END DO
C
IF (LFLAGS(3).EQ.1) THEN
C  Normal incrementation
IF (LFLAGS(1).EQ.1 .OR. LFLAGS(1).EQ.2) THEN
C  *STATIC
  AMATRX(1,1) = AK
  AMATRX(4,4) = AK
  AMATRX(1,4) = -AK
  AMATRX(4,1) = -AK
  IF (LFLAGS(4).NE.0) THEN
    FORCE = AK*(U(4)-U(1))
    DFORCE = AK*(DU(4,1)-DU(1,1))
    SRESID(1) = -DFORCE
    SRESID(4) = DFORCE
    RHS(1,1) = RHS(1,1)-SRESID(1)
    RHS(4,1) = RHS(4,1)-SRESID(4)
    ENERGY(2) = HALF*FORCE*(DU(4,1)-DU(1,1))
    *
    + HALF*DFORCE*(U(4)-U(1))
    *
    + HALF*DFORCE*(DU(4,1)-DU(1,1))
  ELSE
    FORCE = AK*(U(4)-U(1))
    SRESID(1) = -FORCE
    SRESID(4) = FORCE
    RHS(1,1) = RHS(1,1)-SRESID(1)
    RHS(4,1) = RHS(4,1)-SRESID(4)
    DO KDLOAD = 1, NDLOAD
      IF (JDLTYP(KDLOAD,1).EQ.1001) THEN
        RHS(4,1) = RHS(4,1)+ADLMAG(KDLOAD,1)
        ENERGY(8) = ENERGY(8)+(ADLMAG(KDLOAD,1)
        *
        - HALF*DDLMAG(KDLOAD,1))*DU(4,1)
      IF (NRHS.EQ.2) THEN

```

```

C          Riks
          RHS(4,2) = RHS(4,2)+DDLMAG(KDLOAD,1)
          END IF
          END IF
          END DO
          ENERGY(2) = HALF*FORCE*(U(4)-U(1))
          END IF
ELSE IF (LFLAGS(1).EQ.11 .OR. LFLAGS(1).EQ.12) THEN
C          *DYNAMIC
          ALPHA = PARAMS(1)
          BETA = PARAMS(2)
          GAMMA = PARAMS(3)
C          DADU = ONE/(BETA*DTIME**2)
          DVDU = GAMMA/(BETA*DTIME)
C          DO K1 = 1, NDOFEL
          AMATRX(K1,K1) = AM*DADU
          RHS(K1,1) = RHS(K1,1)-AM*A(K1)
          END DO
          AMATRX(1,1) = AMATRX(1,1)+(ONE+ALPHA)*AK
          AMATRX(4,4) = AMATRX(4,4)+(ONE+ALPHA)*AK
          AMATRX(1,4) = AMATRX(1,4)-(ONE+ALPHA)*AK
          AMATRX(4,1) = AMATRX(4,1)-(ONE+ALPHA)*AK
          FORCE = AK*(U(4)-U(1))
          SRESID(1) = -FORCE
          SRESID(4) = FORCE
          RHS(1,1) = RHS(1,1) -
          ((ONE+ALPHA)*SRESID(1)-ALPHA*SVARS(1))
          RHS(4,1) = RHS(4,1) -
          ((ONE+ALPHA)*SRESID(4)-ALPHA*SVARS(4))
          ENERGY(1) = ZERO
          DO K1 = 1, NDOFEL
          SVARS(K1+6) = SVARS(k1)
          SVARS(K1) = SRESID(K1)
          ENERGY(1) = ENERGY(1)+HALF*V(K1)*AM*V(K1)
          END DO
          ENERGY(2) = HALF*FORCE*(U(4)-U(1))
          END IF
ELSE IF (LFLAGS(3).EQ.2) THEN
C          Stiffness matrix
          AMATRX(1,1) = AK

```

```

        AMATRX(4,4) = AK
        AMATRX(1,4) = -AK
        AMATRX(4,1) = -AK
    ELSE IF (LFLAGS(3).EQ.4) THEN
C      Mass matrix
        DO K1 = 1, NDOFEL
            AMATRX(K1,K1) = AM
        END DO
    ELSE IF (LFLAGS(3).EQ.5) THEN
C      Half-increment residual calculation
        ALPHA = PARAMS(1)
        FORCE = AK*(U(4)-U(1))
        SRESID(1) = -FORCE
        SRESID(4) = FORCE
        RHS(1,1) = RHS(1,1)-AM*A(1)-(ONE+ALPHA)*SRESID(1)
        *      + HALF*ALPHA*( SVARS(1)+SVARS(7) )
        RHS(4,1) = RHS(4,1)-AM*A(4)-(ONE+ALPHA)*SRESID(4)
        *      + HALF*ALPHA*( SVARS(4)+SVARS(10) )
    ELSE IF (LFLAGS(3).EQ.6) THEN
C      Initial acceleration calculation
        DO K1 = 1, NDOFEL
            AMATRX(K1,K1) = AM
        END DO
        FORCE = AK*(U(4)-U(1))
        SRESID(1) = -FORCE
        SRESID(4) = FORCE
        RHS(1,1) = RHS(1,1)-SRESID(1)
        RHS(4,1) = RHS(4,1)-SRESID(4)
        ENERGY(1) = ZERO
        DO K1 = 1, NDOFEL
            SVARS(K1) = SRESID(K1)
            ENERGY(1) = ENERGY(1)+HALF*V(K1)*AM*V(K1)
        END DO
        ENERGY(2) = HALF*FORCE*(U(4)-U(1))
    ELSE IF (LFLAGS(3).EQ.100) THEN
C      Output for perturbations
        IF (LFLAGS(1).EQ.1 .OR. LFLAGS(1).EQ.2) THEN
            *STATIC
            FORCE = AK*(U(4)-U(1))
            DFORCE = AK*(DU(4,1)-DU(1,1))
            SRESID(1) = -DFORCE
            SRESID(4) = DFORCE

```

```
RHS(1,1) = RHS(1,1)-SRESID(1)
RHS(4,1) = RHS(4,1)-SRESID(4)
ENERGY(2) = HALF*FORCE*(DU(4,1)-DU(1,1))
*          + HALF*DFORCE*(U(4)-U(1))
*          + HALF*DFORCE*(DU(4,1)-DU(1,1))
DO KVAR = 1, NSVARS
    SVARS(KVAR) = ZERO
END DO
SVARS(1) = RHS(1,1)
SVARS(4) = RHS(4,1)
ELSE IF (LFLAGS(1).EQ.41) THEN
    C
        *FREQUENCY
        DO KRHS = 1, NRHS
            DFORCE = AK*(DU(4,KRHS)-DU(1,KRHS))
            SRESID(1) = -DFORCE
            SRESID(4) = DFORCE
            RHS(1,KRHS) = RHS(1,KRHS)-SRESID(1)
            RHS(4,KRHS) = RHS(4,KRHS)-SRESID(4)
        END DO
        DO KVAR = 1, NSVARS
            SVARS(KVAR) = ZERO
        END DO
        SVARS(1) = RHS(1,1)
        SVARS(4) = RHS(4,1)
    END IF
END IF
C
RETURN
END
```

## 1.1.25 UELMAT: User subroutine to define an element with access to Abaqus materials.

**Product:** Abaqus/Standard

*WARNING: This feature is intended for advanced users only. Its use in all but the simplest test examples will require considerable coding by the user/developer.*

*“User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual, should be read before proceeding.*

---

### References

- “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual
- \*UEL PROPERTY
- \*USER ELEMENT
- “Accessing Abaqus materials,” Section 2.1.17
- “Accessing Abaqus thermal materials,” Section 2.1.18

---

### Overview

User subroutine **UELMAT**:

- will be called for each element that is of a general user-defined element type (i.e., not defined by a linear stiffness or mass matrix read either directly or from results file data) each time element calculations are required;
- (or subroutines called by user subroutine **UELMAT**) must perform all of the calculations for the element, appropriate to the current activity in the analysis;
- can access some of the Abaqus materials through utility routines **MATERIAL\_LIB\_MECH** and **MATERIAL\_LIB\_HT**;
- is available for a subset of the procedures supported for user subroutine **UEL** (see “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual); and
- is available for plane stress and three-dimensional element types in a stress/displacement analysis and for two-dimensional and three-dimensional element types in a heat transfer analysis (see “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

---

### User subroutine interface

```
SUBROUTINE UELMAT (RHS ,AMATRX ,SVARS ,ENERGY ,NDOFEL ,NRHS ,NSVARS ,
1 PROPS ,NPROPS ,COORDS ,MCRD ,NNODE ,U ,DU ,V ,A ,JTYPE ,TIME ,DTIME ,
2 KSTEP ,KINC ,JELEM ,PARAMS ,NDLOAD ,JDLTYP ,ADLMAG ,PREDEF ,NPREFD ,
3 LFLAGS ,MLVARX ,DDLMAG ,MDLOAD ,PNEWDT ,JPROPS ,NJPROP ,PERIOD ,
4 MATERIALLIB )
```

```

C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION RHS (MLVARX,*),AMATRX (NDOFEL,NDOFEL),PROPS (*),
1 SVARS (*),ENERGY (8),COORDS (MCRD,NNODE),U (NDOFEL),
2 DU (MLVARX,*),V (NDOFEL),A (NDOFEL),TIME (2),PARAMS (*),
3 JDLTYP (MDLOAD,*),ADLMAG (MDLOAD,*),DDLMAG (MDLOAD,*),
4 PREDEF (2,NPREDF,NNODE),LFLAGS (*),JPROPS (*)

```

*user coding to define **RHS**, **AMATRX**, **SVARS**, **ENERGY**, and **PNEWDT***

```

RETURN
END

```

## Variables to be defined

---

These arrays depend on the value of the **LFLAGS** array.

### RHS

An array containing the contributions of this element to the right-hand-side vectors of the overall system of equations. For most nonlinear analysis procedures, **NRHS**=1 and **RHS** should contain the residual vector. The exception is the modified Riks static procedure (“Static stress analysis,” Section 6.2.2 of the Abaqus Analysis User’s Manual), for which **NRHS**=2 and the first column in **RHS** should contain the residual vector and the second column should contain the increments of external load on the element. **RHS (K1, K2)** is the entry for the **K1**th degree of freedom of the element in the **K2**th right-hand-side vector.

### AMATRX

An array containing the contribution of this element to the Jacobian (stiffness) or other matrix of the overall system of equations. The particular matrix required at any time depends on the entries in the **LFLAGS** array (see below).

All nonzero entries in **AMATRX** should be defined, even if the matrix is symmetric. If you do not specify that the matrix is unsymmetric when you define the user element, Abaqus/Standard will use the symmetric matrix defined by  $\frac{1}{2}([A] + [A]^T)$ , where  $[A]$  is the matrix defined as **AMATRX** in this subroutine. If you specify that the matrix is unsymmetric when you define the user element, Abaqus/Standard will use **AMATRX** directly.

### SVARS

An array containing the values of the solution-dependent state variables associated with this element. The number of such variables is **NSVARS** (see below). You define the meaning of these variables.

For general nonlinear steps this array is passed into **UELMAT** containing the values of these variables at the start of the current increment. They should be updated to be the values at the end

of the increment, unless the procedure during which **UELMAT** is being called does not require such an update; this requirement depends on the entries in the **LFLAGS** array (see below). For linear perturbation steps this array is passed into **UELMAT** containing the values of these variables in the base state. They should be returned containing perturbation values if you wish to output such quantities.

When **KINC** is equal to zero, the call to **UELMAT** is made for zero increment output (see “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). In this case the values returned will be used only for output purposes and are not updated permanently.

### **ENERGY**

For general nonlinear steps array **ENERGY** contains the values of the energy quantities associated with the element. The values in this array when **UELMAT** is called are the element energy quantities at the start of the current increment. They should be updated to the values at the end of the current increment. For linear perturbation steps the array is passed into **UELMAT** containing the energy in the base state. They should be returned containing perturbation values if you wish to output such quantities. The entries in the array are as follows:

<b>ENERGY (1)</b>	Kinetic energy.
<b>ENERGY (2)</b>	Elastic strain energy.
<b>ENERGY (3)</b>	Creep dissipation.
<b>ENERGY (4)</b>	Plastic dissipation.
<b>ENERGY (5)</b>	Viscous dissipation.
<b>ENERGY (6)</b>	“Artificial strain energy” associated with such effects as artificial stiffness introduced to control hourgassing or other singular modes in the element.
<b>ENERGY (7)</b>	Electrostatic energy.
<b>ENERGY (8)</b>	Incremental work done by loads applied within the user element.

When **KINC** is equal to zero, the call to **UELMAT** is made for zero increment output (see “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). In this case the energy values returned will be used only for output purposes and are not updated permanently.

### **Variable that can be updated**

---

#### **PNEWDT**

Ratio of suggested new time increment to the time increment currently being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen). It is useful only during equilibrium iterations with the normal time incrementation, as indicated by **LFLAGS (3)=1**. During a severe discontinuity iteration (such as contact changes), **PNEWDT** is ignored unless CONVERT SDI=YES is specified for this step. The usage of **PNEWDT** is discussed below.

**PNEWDT** is set to a large value before each call to **UELMAT**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

## Variables passed in for information

---

### Arrays:

#### **PROPS**

A floating point array containing the **NPROPS** real property values defined for use with this element. **NPROPS** is the user-specified number of real property values. See “Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual.

#### **JPROPS**

An integer array containing the **NJPROP** integer property values defined for use with this element. **NJPROP** is the user-specified number of integer property values. See “Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual.

#### **COORDS**

An array containing the original coordinates of the nodes of the element. **COORDS (K1 , K2)** is the **K1**th coordinate of the **K2**th node of the element.

#### **U, DU, V, A**

Arrays containing the current estimates of the basic solution variables (displacements, rotations, temperatures, depending on the degree of freedom) at the nodes of the element at the end of the current increment. Values are provided as follows:

**U (K1)** Total values of the variables. If this is a linear perturbation step, it is the value in the base state.

**DU (K1 , KRHS)** Incremental values of the variables for the current increment for right-hand-side **KRHS**. If this is an eigenvalue extraction step, this is the eigenvector magnitude for eigenvector **KRHS**. For steady-state dynamics **KRHS** = 1 denotes real components of perturbation displacement and **KRHS** = 2 denotes imaginary components of perturbation displacement.

<b>V (K1)</b>	Time rate of change of the variables (velocities, rates of rotation). Defined for implicit dynamics only ( <b>LFLAGS (1)</b> = 11 or 12).
<b>A (K1)</b>	Accelerations of the variables. Defined for implicit dynamics only ( <b>LFLAGS (1)</b> = 11 or 12).

**JDLTYP**

An array containing the integers used to define distributed load types for the element. Loads of type  $U_n$  are identified by the integer value  $n$  in **JDLTYP**; loads of type  $U_nNU$  are identified by the negative integer value  $-n$  in **JDLTYP**. **JDLTYP (K1, K2)** is the identifier of the **K1**th distributed load in the **K2**th load case. For general nonlinear steps **K2** is always 1.

**ADLMAG**

For general nonlinear steps **ADLMAG (K1, 1)** is the total load magnitude of the **K1**th distributed load at the end of the current increment for distributed loads of type  $U_n$ . For distributed loads of type  $U_nNU$ , the load magnitude is defined in **UELMAT**; therefore, the corresponding entries in **ADLMAG** are zero. For linear perturbation steps **ADLMAG (K1, 1)** contains the total load magnitude of the **K1**th distributed load of type  $U_n$  applied in the base state. Base state loading of type  $U_nNU$  must be dealt with inside **UELMAT**. **ADLMAG (K1, 2)**, **ADLMAG (K1, 3)**, etc. are currently not used.

**DDLMAG**

For general nonlinear steps **DDLMAG** contains the increments in the magnitudes of the distributed loads that are currently active on this element for distributed loads of type  $U_n$ . **DDLMAG (K1, 1)** is the increment of magnitude of the load for the current time increment. The increment of load magnitude is needed to compute the external work contribution. For distributed loads of type  $U_nNU$  the load magnitude is defined in **UELMAT**; therefore, the corresponding entries in **DDLMAG** are zero. For linear perturbation steps **DDLMAG (K1, K2)** contains the perturbation in the magnitudes of the distributed loads that are currently active on this element for distributed loads of type  $U_n$ . **K1** denotes the **K1**th perturbation load active on the element. **K2** is always 1, except for steady-state dynamics, where **K2=1** for real loads and **K2=2** for imaginary loads. Perturbation loads of type  $U_nNU$  must be dealt with inside **UELMAT**.

**PREDEF**

An array containing the values of predefined field variables, such as temperature in an uncoupled stress/displacement analysis, at the nodes of the element (“Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual).

The first index of the array, **K1**, is either 1 or 2, with 1 indicating the value of the field variable at the end of the increment and 2 indicating the increment in the field variable. The second index, **K2**, indicates the variable: the temperature corresponds to index 1, and the predefined field variables correspond to indices 2 and above. In cases where temperature is not defined, the predefined field variables begin with index 1. The third index, **K3**, indicates the local node number on the element.

<b>PREDEF (K1 , 1 , K3)</b>	Temperature.
<b>PREDEF (K1 , 2 , K3)</b>	First predefined field variable.
<b>PREDEF (K1 , 3 , K3)</b>	Second predefined field variable.
Etc.	Any other predefined field variable.
<b>PREDEF (K1 , K2 , K3)</b>	Total or incremental value of the <b>K2</b> th predefined field variable at the <b>K3</b> th node of the element.
<b>PREDEF (1 , K2 , K3)</b>	Values of the variables at the end of the current increment.
<b>PREDEF (2 , K2 , K3)</b>	Incremental values corresponding to the current time increment.

**PARAMS**

An array containing the parameters associated with the solution procedure. The entries in this array depend on the solution procedure currently being used when **UELMAT** is called, as indicated by the entries in the **LFLAGS** array (see below).

For implicit dynamics (**LFLAGS (1) = 11** or **12**) **PARAMS** contains the integration operator values, as:

<b>PARAMS (1)</b>	$\alpha$
<b>PARAMS (2)</b>	$\beta$
<b>PARAMS (3)</b>	$\gamma$

**LFLAGS**

An array containing the flags that define the current solution procedure and requirements for element calculations. Detailed requirements for the various Abaqus/Standard procedures are defined earlier in this section.

<b>LFLAGS (1)</b>	Defines the procedure type. See “Results file output format,” Section 5.1.2 of the Abaqus Analysis User’s Manual, for the key used for each procedure.
<b>LFLAGS (2)=0</b>	Small-displacement analysis.
<b>LFLAGS (2)=1</b>	Large-displacement analysis (nonlinear geometric effects included in the step; see “General and linear perturbation procedures,” Section 6.1.2 of the Abaqus Analysis User’s Manual).
<b>LFLAGS (3)=1</b>	Normal implicit time incrementation procedure. User subroutine <b>UELMAT</b> must define the residual vector in <b>RHS</b> and the Jacobian matrix in <b>AMATRX</b> .
<b>LFLAGS (3)=2</b>	Define the current stiffness matrix ( <b>AMATRX</b> = $K^{NM} = -\partial F^N / \partial u^M$ or $-\partial G^N / \partial u^M$ ) only.

<b>LFLAGS (3)=3</b>	Define the current damping matrix ( <b>AMATRX</b> = $C^{NM} = -\partial F^N / \partial u^M$ or $-\partial G^N / \partial u^M$ ) only.
<b>LFLAGS (3)=4</b>	Define the current mass matrix ( <b>AMATRX</b> = $M^{NM} = -\partial F^N / \partial \ddot{u}^M$ ) only. Abaqus/Standard always requests an initial mass matrix at the start of the analysis.
<b>LFLAGS (3)=5</b>	Define the current residual or load vector ( <b>RHS</b> = $F^N$ ) only.
<b>LFLAGS (3)=6</b>	Define the current mass matrix and the residual vector for the initial acceleration calculation (or the calculation of accelerations after impact).
<b>LFLAGS (3)=100</b>	Define perturbation quantities for output.
<b>LFLAGS (4)=0</b>	The step is a general step.
<b>LFLAGS (4)=1</b>	The step is a linear perturbation step.
<b>LFLAGS (5)=0</b>	The current approximations to $u^M$ , etc. were based on Newton corrections.
<b>LFLAGS (5)=1</b>	The current approximations were found by extrapolation from the previous increment.

**TIME (1)**

Current value of step time.

**TIME (2)**

Current value of total time.

**Scalar parameters:****DTIME**

Time increment.

**PERIOD**

Time period of the current step.

**NDOFEL**

Number of degrees of freedom in the element.

**MLVARX**

Dimensioning parameter used when several displacement or right-hand-side vectors are used.

**NRHS**

Number of load vectors. **NRHS** is 1 in most nonlinear problems: it is 2 for the modified Riks static procedure (“Static stress analysis,” Section 6.2.2 of the Abaqus Analysis User’s Manual), and it is greater than 1 in some linear analysis procedures and during substructure generation.

**NSVARS**

User-defined number of solution-dependent state variables associated with the element (“Defining the number of solution-dependent variables that must be stored within the element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**NPROPS**

User-defined number of real property values associated with the element (“Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**NJPROP**

User-defined number of integer property values associated with the element (“Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**MCRD**

**MCRD** is defined as the maximum of the user-defined maximum number of coordinates needed at any node point (“Defining the maximum number of coordinates needed at any nodal point” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual) and the value of the largest active degree of freedom of the user element that is less than or equal to 3. For example, if you specify that the maximum number of coordinates is 1 and the active degrees of freedom of the user element are 2, 3, and 6, **MCRD** will be 3. If you specify that the maximum number of coordinates is 2 and the active degrees of freedom of the user element are 11 and 12, **MCRD** will be 2.

**NNODE**

User-defined number of nodes on the element (“Defining the number of nodes associated with the element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**JTYPE**

Integer defining the element type. This is the user-defined integer value  $n$  in element type  $Un$  (“Assigning an element type key to a user-defined element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**KSTEP**

Current step number.

**KINC**

Current increment number.

**JELEM**

User-assigned element number.

**NDLOAD**

Identification number of the distributed load or flux currently active on this element.

**MDLOAD**

Total number of distributed loads and/or fluxes defined on this element.

**NPREFD**

Number of predefined field variables, including temperature. For user elements Abaqus/Standard uses one value for each field variable per node.

**MATERIALLIB**

A variable that must be passed to the utility routines performing material point computations.

**UELMAT conventions**

---

The solution variables (displacement, velocity, etc.) are arranged on a node/degree of freedom basis. The degrees of freedom of the first node are first, followed by the degrees of freedom of the second node, etc.

**Usage with general nonlinear procedures**

---

The values of  $u^N$  (and, in direct-integration dynamic steps,  $\dot{u}^N$  and  $\ddot{u}^N$ ) enter user subroutine **UELMAT** as their latest approximations at the end of the time increment; that is, at time  $t + \Delta t$ .

The values of  $H^\alpha$  enter the subroutine as their values at the beginning of the time increment; that is, at time  $t$ . It is your responsibility to define suitable time integration schemes to update  $H^\alpha$ . To ensure accurate, stable integration of internal state variables, you can control the time incrementation via **PNEWDT**.

The values of  $p^\beta$  enter the subroutine as the values of the total load magnitude for the  $\beta$ th distributed load at the end of the increment. Increments in the load magnitudes are also available.

In the following descriptions of the user element's requirements, it will be assumed that **LFLAGS (3)=1** unless otherwise stated.

**Static analysis (**LFLAGS (1)=1, 2**)**

- $F^N = F^N(u^M, H^\alpha, p^\beta, t)$ .
- Automatic convergence checks are applied to the force residuals corresponding to degrees of freedom 1–7.
- You must define **AMATRX** =  $K^{NM} = -\partial F^N / \partial u^M$  and **RHS** =  $F^N$  and update the state variables,  $H^\alpha$ .

**Direct-integration dynamic analysis (**LFLAGS (1)=11, 12**)**

- Automatic convergence checks are applied to the force residuals corresponding to degrees of freedom 1–7.
- **LFLAGS (3)=1**: Normal time increment. Either the Hilber-Hughes-Taylor or the backward Euler time integration scheme will be used. With  $\alpha$  set to zero for the backward Euler, both schemes imply

$$F^N = -M^{NM}\ddot{u}_{t+\Delta t} + (1 + \alpha)G^N - \alpha G_t^N,$$

where  $M^{NM} = M^{NM}(u^M, \dot{u}^M, H^\alpha, p^\beta, t, \dots)$  and  $G^N = G^N(u^M, \dot{u}^M, H^\alpha, p^\beta, t, \dots)$ ; that is, the highest time derivative of  $u^M$  in  $M^{NM}$  and  $G^N$  is  $\dot{u}^M$ , so that

$$-\frac{\partial F^N}{\partial \dot{u}^M_{t+\Delta t}} = M^{NM}.$$

Therefore, you must store  $G_t^N$  as an internal state vector. If half-increment residual calculations are required, you must also store  $G_{t^-}^N$  as an internal state vector, where  $t^-$  indicates the time at the beginning of the previous increment. For  $\alpha = 0$ ,  $F^N = -M^{NM}\ddot{u}_{t+\Delta t} + G_{t+\Delta t}^N$  and  $G_t^N$  is not needed. You must define **AMATRX** =  $M^{NM}(d\ddot{u}/du) + (1 + \alpha)C^{NM}(d\dot{u}/du) + (1 + \alpha)K^{NM}$ , where  $C^{NM} = -\partial G^N_{t+\Delta t}/\partial \dot{u}^M$  and  $K^{NM} = -\partial G^N_{t+\Delta t}/\partial u^M$ . **RHS** =  $F^N$  must also be defined and the state variables,  $H^\alpha$ , updated. Although the value of  $\alpha$  given in the dynamic step definition is passed into **UELMAT**, the value of  $\alpha$  can vary from element to element. For example,  $\alpha$  can be set to zero for some elements in the model where numerical dissipation is not desired.

- **LFLAGS (3)=5:** Half-increment residual ( $F_{1/2}^N$ ) calculation. Abaqus/Standard will adjust the time increment so that  $\max |F_{1/2}^N| < tolerance$  (where *tolerance* is specified in the dynamic step definition). The half-increment residual is defined as

$$F_{1/2}^N = -M^{NM}\ddot{u}_{t+\Delta t/2} + (1 + \alpha)G_{t+\Delta t/2}^N - \frac{\alpha}{2}(G_t^N + G_{t^-}^N),$$

where  $t^-$  indicates the time at the beginning of the previous increment ( $\alpha$  is a parameter of the Hilber-Hughes-Taylor time integration operator and will be set to zero if the backward Euler time integration operator is used). You must define **RHS** =  $F_{1/2}^N$ . To evaluate  $M^{NM}$  and  $G_{t+\Delta t/2}^N$ , you must calculate  $H^\alpha_{t+\Delta t/2}$ . These half-increment values will not be saved. **DTIME** will still contain  $\Delta t$  (not  $\Delta t/2$ ). The values contained in **U**, **V**, **A**, and **DU** are half-increment values.

- **LFLAGS (3)=4:** Velocity jump calculation. Abaqus/Standard solves  $-M^{NM}\Delta\dot{u}^M = 0$  for  $\Delta\dot{u}^M$ , so you must define **AMATRX** =  $M^{NM}$ .
- **LFLAGS (3)=6:** Initial acceleration calculation. Abaqus/Standard solves  $-M^{NM}\ddot{u}^M + G^N = 0$  for  $\ddot{u}^M$ , so you must define **AMATRX** =  $M^{NM}$  and **RHS** =  $G^N$ .

### Quasi-static analysis (**LFLAGS (1)=21**)

- The requirements are identical to those of static analysis.

### Steady-state heat transfer analysis (**LFLAGS (1)=31**)

- The requirements are identical to those of static analysis, except that the automatic convergence checks are applied to the heat flux residuals corresponding to degrees of freedom 11, 12, ...

### Transient heat transfer analysis ( $\theta_{max}$ ) (**LFLAGS (1)=32, 33**)

- Automatic convergence checks are applied to the heat flux residuals corresponding to degrees of freedom 11, 12, ...

- The backward difference scheme is always used for time integration; that is, Abaqus/Standard assumes that  $\dot{u}_{t+\Delta t} = \Delta u / \Delta t$ , where  $\Delta u = u_{t+\Delta t} - u_t$  and so  $d\dot{u}/du = 1/\Delta t$  always. For degrees of freedom 11, 12, ...,  $\max |\Delta u|$  will be compared against the user-prescribed maximum allowable nodal temperature change in an increment,  $\Delta\theta_{max}$ , for controlling the time integration accuracy.
- You need to define  $\text{AMATRX} = K^{NM} + (1/\Delta t) C^{NM}$ , where  $C^{NM}$  is the heat capacity matrix and  $\text{RHS} = F^N$ , and must update the state variables,  $H^\alpha$ .

## Usage with linear perturbation procedures

---

“General and linear perturbation procedures,” Section 6.1.2 of the Abaqus Analysis User’s Manual, describes the linear perturbation capabilities in Abaqus/Standard. Here, base state values of variables will be denoted by  $u^M$ ,  $H^\alpha$ , etc. Perturbation values will be denoted by  $\tilde{u}^M$ ,  $\tilde{H}^\alpha$ , etc.

Abaqus/Standard will not call user subroutine **UELMAT** for the following procedures: eigenvalue buckling prediction, response spectrum, transient modal dynamic, steady-state dynamic (modal and direct), and random response.

### Static analysis (**LFLAGS (1)=1, 2**)

- Abaqus/Standard will solve  $K^{NM} \tilde{u}^M = \tilde{P}^N$  for  $\tilde{u}^M$ , where  $K^{NM}$  is the base state stiffness matrix and the perturbation load vector,  $\tilde{P}^N$ , is a linear function of the perturbation loads,  $\tilde{p}$ ; that is,  $\tilde{P}^N = (\partial F / \partial \tilde{p}) \tilde{p}$ .
- LFLAGS (3)=1:** You must define  $\text{AMATRX} = K^{NM}$  and  $\text{RHS} = \tilde{P}^N$ .
- LFLAGS (3)=100:** You must compute perturbations of the internal variables,  $\tilde{H}^\alpha$ , and define  $\text{RHS} = \tilde{P}^N - K^{NM} \tilde{u}^M$  for output purposes.

### Eigenfrequency extraction analysis (**LFLAGS (1)=41**)

- $F^N = -M^{NM} \ddot{\tilde{u}} + G^N(u^M + \tilde{u}^M, \dots) = -M^{NM} \ddot{\tilde{u}} + (\partial G^N / \partial u^M) \tilde{u}^M$ .
- Abaqus/Standard will solve  $K^{NM} \phi_i^M = \omega_i^2 M^{NM} \phi_i^M$  for  $\phi_i^N$  and  $\omega_i$ , where  $K^{NM} = -\partial F^N / \partial u^M$  is the base state stiffness matrix and  $M^{NM} = -\partial F^{NM} / \partial \ddot{u}^M$  is the base state mass matrix.
- LFLAGS (3)=2:** Define  $\text{AMATRX} = K^{NM}$ .
- LFLAGS (3)=4:** Define  $\text{AMATRX} = M^{NM}$ .

### Example: Structural user element with Abaqus isotropic linearly elastic material

---

Both a structural and a heat transfer user element have been created to demonstrate the usage of subroutine **UELMAT**. These user-defined elements are applied in a number of analyses. The following excerpt illustrates how the linearly elastic isotropic material available in Abaqus can be accessed from user subroutine **UELMAT**:

```
...
*USER ELEMENT, TYPE=U1, NODES=4, COORDINATES=2, VAR=16,
INTEGRATION=4, TENSOR=PSTRAIN
```

```

1,2
*ELEMENT, TYPE=U1, ELSET=SOLID
1, 1,2,3,4
...
*UEL PROPERTY, ELSET=SOLID, MATERIAL=MAT
...
*MATERIAL, NAME=MAT
*ELASTIC
7.00E+010, 0.33

```

The user element defined above is a 4-node, fully integrated plane strain element, similar to the Abaqus CPE4 element.

The next excerpt shows the listing of the user subroutine. Inside the subroutine, a loop over the integration points is performed. For each integration point the utility routine **MATERIAL\_LIB\_MECH** is called, which returns stress and Jacobian at the integration point. These quantities are used to compute the right-hand-side vector and the element Jacobian.

```

c*****
      subroutine uelmat(rhs,amatrix,svars,energy,ndofel,nrhs,
1      nsvars,props,nprops,coords,mcrd,nnode,u,du,
2      v,a,jtype,time,dtime,kstep,kinc,jelem,params,
3      ndload,jdltyp,adlmag,predef,npredf,lfflags,mlvarx,
4      ddlmag,mdload,pnewdt,jprops,njpro,period,
5      materiallib)
c
      include 'aba_param.inc'
c
      dimension rhs(mlvarx,*), amatrix(ndofel, ndofel), props(*),
1      svars(*), energy(*), coords(mcrd, nnode), u(ndofel),
2      du(mlvarx,*), v(ndofel), a(ndofel), time(2), params(*),
3      jdltyp(mdload,*), adlmag(mdload,*), ddlmag(mdload,*),
4      predef(2, npredf, nnode), lfflags(*), jprops(*)
      parameter (zero=0.d0, dmone=-1.0d0, one=1.d0, four=4.0d0,
1      fourth=0.25d0,gaussCoord=0.577350269d0)
      parameter (ndim=2, ndof=2, nshr=1, nnodemax=4,
1      ntens=4, ninpt=4, nsint=4)
c
c      ndim ... number of spatial dimensions
c      ndof ... number of degrees of freedom per node
c      nshr ... number of shear stress component
c      ntens ... total number of stress tensor components
c                  (=ndi+nshr)

```

```

c      ninpt ... number of integration points
c      nsvint... number of state variables per integration pt
c                  (strain)
c
c      dimension stiff(ndof*nnodemax,ndof*nnodemax),
1 force(ndof*nnodemax), shape(nnodemax), dshape(ndim,nnodemax),
2 xjac(ndim,ndim), xjaci(ndim,ndim), bmat(nnodemax*ndim),
3 statevLocal(nsvint), stress(ntens), ddsdde(ntens, ntens),
4 stran(ntens), dstran(ntens), wght(ninpt)
c
c      dimension predef_loc(npredf),dpredef_loc(npredf),
1      defGrad(3,3),utmp(3),xdu(3),stiff_p(3,3),force_p(3)
dimension coord24(2,4),coords_ip(3)
data coord24 /dmone, dmone,
2           one, dmone,
3           one,   one,
4           dmone, one/
c
data wght /one, one, one, one/
c*****
c
c      U1 = first-order, plane strain, full integration
c
c      State variables: each integration point has nsvint SDVs
c
c      isvinc=(npt-1)*nsvint    ... integration point counter
c      statev(1+isvinc        ) ... strain
c
c*****
if (lflags(3).eq.4) then
  do i=1, ndofel
    do j=1, ndofel
      amatrx(i,j) = zero
    end do
    amatrx(i,i) = one
  end do
  goto 999
end if
c
c      PRELIMINARIES
c

```

```

pnewdtLocal = pnewdt
if(jtype .ne. 1) then
    write(7,*)'Incorrect element type'
    call xit
endif
if(nsvars .lt. ninpt*nsvint) then
    write(7,*)'Increase the number of SDVs to', ninpt*nsvint
    call xit
endif
thickness = 0.1d0

c
c      INITIALIZE RHS AND LHS
c
do k1=1, ndof*nnode
    rhs(k1, 1)= zero
    do k2=1, ndof*nnode
        amatrix(k1, k2)= zero
    end do
end do

c
c      LOOP OVER INTEGRATION POINTS
c
do kintk = 1, ninpt
c
c      EVALUATE SHAPE FUNCTIONS AND THEIR DERIVATIVES
c
c      determine (g,h)
c
g = coord24(1,kintk)*gaussCoord
h = coord24(2,kintk)*gaussCoord
c
c      shape functions
shape(1) = (one - g)*(one - h)/four;
shape(2) = (one + g)*(one - h)/four;
shape(3) = (one + g)*(one + h)/four;
shape(4) = (one - g)*(one + h)/four;
c
c      derivative d(Ni)/d(g)
dshape(1,1) = -(one - h)/four;
dshape(1,2) = (one - h)/four;
dshape(1,3) = (one + h)/four;
dshape(1,4) = -(one + h)/four;

```

```

c
c      derivative d(Ni)/d(h)
dshape(2,1) = -(one - g)/four;
dshape(2,2) = -(one + g)/four;
dshape(2,3) = (one + g)/four;
dshape(2,4) = (one - g)/four;
c
c      compute coordinates at the integration point
c
do k1=1, 3
    coords_ip(k1) = zero
end do
do k1=1,nnode
    do k2=1,mcrd
        coords_ip(k2)=coords_ip(k2)+shape(k1)*coords(k2,k1)
    end do
end do
c
c      INTERPOLATE FIELD VARIABLES
c
if(npredf.gt.0) then

    do k1=1,npredf
        predef_loc(k1) = zero
        dpredef_loc(k1) = zero
        do k2=1,nnode
            predef_loc(k1) =
&             predef_loc(k1) +
&             (predef(1,k1,k2)-predef(2,k1,k2))*shape(k2)
            dpredef_loc(k1) =
&             dpredef_loc(k1)+predef(2,k1,k2)*shape(k2)
        end do
    end do
end if
c
c      FORM B-MATRIX
c
djac = one
c
do i = 1, ndim
    do j = 1, ndim
        xjac(i,j) = zero

```

```

        xjaci(i,j) = zero
    end do
end do

c
do inod= 1, nnnode
    do idim = 1, ndim
        do jdim = 1, ndim
            xjac(jdim,idim) = xjac(jdim,idim) +
1           dshape(jdim,inod)*coords(idim,inod)
        end do
    end do
end do
djac = xjac(1,1)*xjac(2,2) - xjac(1,2)*xjac(2,1)
if (djac .gt. zero) then
    ! jacobian is positive - o.k.
    xjaci(1,1) = xjac(2,2)/djac
    xjaci(2,2) = xjac(1,1)/djac
    xjaci(1,2) = -xjac(1,2)/djac
    xjaci(2,1) = -xjac(2,1)/djac
else
    ! negative or zero jacobian
    write(7,*) 'WARNING: element',jelem,'has neg.
1          Jacobian'
    pnewdt = fourth
endif

if (pnewdt .lt. pnewdtLocal) pnewdtLocal = pnewdt

c
do i = 1, nnnode*ndim
    bmat(i) = zero
end do

do inod = 1, nnnode
    do ider = 1, ndim
        do idim = 1, ndim
            irow = idim + (inod - 1)*ndim
            bmat(irow) = bmat(irow) +
1           xjaci(idim,ider)*dshape(ider,inod)
        end do
    end do
end do

```

```

c
c      CALCULATE INCREMENTAL STRAINS
c
do i = 1, ntens
  dstran(i) = zero
end do
!
! set deformation gradient to Identity matrix
do k1=1,3
  do k2=1,3
    defGrad(k1,k2) = zero
  end do
  defGrad(k1,k1) = one
end do

c
c      COMPUTE INCREMENTAL STRAINS
c
do nodi = 1, nnodes

  incr_row = (nodi - 1)*ndof

  do i = 1, ndof
    xdu(i)= du(i + incr_row,1)
    utmp(i) = u(i + incr_row)
  end do

  dNidx = bmat(1 + (nodi-1)*ndim)
  dNidy = bmat(2 + (nodi-1)*ndim)

  dstran(1) = dstran(1) + dNidx*xdu(1)
  dstran(2) = dstran(2) + dNidy*xdu(2)
  dstran(4) = dstran(4) +
  1      dNidy*xdu(1) +
  2      dNidx*xdu(2)

c      deformation gradient

  defGrad(1,1) = defGrad(1,1) + dNidx*utmp(1)
  defGrad(1,2) = defGrad(1,2) + dNidy*utmp(1)
  defGrad(2,1) = defGrad(2,1) + dNidx*utmp(2)
  defGrad(2,2) = defGrad(2,2) + dNidy*utmp(2)

```

```
    end do

c
c      CALL CONSTITUTIVE ROUTINE
c

      isvinc= (kintk-1)*nsvint ! integration point increment
c
c      prepare arrays for entry into material routines
c

      do i = 1, nsvint
        statevLocal(i)=svars(i+isvinc)
      end do

c
c      state variables
c

      do k1=1,ntens
        stran(k1) = statevLocal(k1)
        stress(k1) = zero
      end do

c
      do i=1, ntens
        do j=1, ntens
          ddsdde(i,j) = zero
        end do
        ddsdde(i,j) = one
      enddo

c
c      compute characteristic element length
c

      celent = sqrt(djac*dble(ninpt))
      dpmat = djac*thickness

c
      dvdv0 = one
      call material_lib_mech(materiallib,stress,ddsdde,
1           stran,dstran,kintk,dvdv0,dpmat,defGrad,
2           predef_loc,dpredef_loc,npredf,celent,coords_ip)

c
      do k1=1,ntens
        statevLocal(k1) = stran(k1) + dstran(k1)
      end do

c
```

```

      isvinc= (kintk-1)*nsvint ! integration point increment
c
c      update element state variables
c
      do i = 1, nsvint
         svars(i+isvinc)=statevLocal(i)
      end do
c
c      form stiffness matrix and internal force vector
c
      dNjdx = zero
      dNjdy = zero
      do i = 1, ndof*nnode
         force(i) = zero
         do j = 1, ndof*nnode
            stiff(j,i) = zero
         end do
      end do

      dvol= wght(kintk)*djac
      do nodj = 1, nnode

         incr_col = (nodj - 1)*ndof

         dNjdx = bmat(1+(nodj-1)*ndim)
         dNjdy = bmat(2+(nodj-1)*ndim)
         force_p(1) = dNjdx*stress(1) + dNjdy*stress(4)
         force_p(2) = dNjdy*stress(2) + dNjdx*stress(4)
         do jdof = 1, ndof

            jcol = jdof + incr_col

            force(jcol) = force(jcol) +
                           force_p(jdof)*dvol
         end do

         do nodi = 1, nnode

            incr_row = (nodi -1)*ndof

            dNidx = bmat(1+(nodi-1)*ndim)

```

```

dNidy = bmat(2+(nodi-1)*ndim)
stiff_p(1,1) = dNidx*ddsdde(1,1)*dNjdx
&           + dNidy*ddsdde(4,4)*dNjdy
&           + dNidx*ddsdde(1,4)*dNjdy
&           + dNidy*ddsdde(4,1)*dNjdx

stiff_p(1,2) = dNidx*ddsdde(1,2)*dNjdy
&           + dNidy*ddsdde(4,4)*dNjdx
&           + dNidx*ddsdde(1,4)*dNjdx
&           + dNidy*ddsdde(4,2)*dNjdy

stiff_p(2,1) = dNidy*ddsdde(2,1)*dNjdx
&           + dNidx*ddsdde(4,4)*dNjdy
&           + dNidy*ddsdde(2,4)*dNjdy
&           + dNidx*ddsdde(4,1)*dNjdx

stiff_p(2,2) = dNidy*ddsdde(2,2)*dNjdy
&           + dNidx*ddsdde(4,4)*dNjdx
&           + dNidy*ddsdde(2,4)*dNjdx
&           + dNidx*ddsdde(4,2)*dNjdy

do jdof = 1, ndof
    icol = jdof + incr_col
    do idof = 1, ndof
        irow = idof + incr_row
        stiff(irow,icol) = stiff(irow,icol) +
&                      stiff_p(idof,jdof)*dvol
    end do
    end do
    end do
end do

c
c      assemble rhs and lhs
c
do k1=1, ndof*nnode
    rhs(k1, 1) = rhs(k1, 1) - force(k1)
    do k2=1, ndof*nnode
        amatr(x(k1, k2) = amatr(x(k1, k2) + stiff(k1,k2)
    end do
    end do
end do      ! end loop on material integration points

```

```
pnewdt = pnewdtLocal  
c  
999  continue  
c  
      return  
end
```



## 1.1.26 UEXPAN: User subroutine to define incremental thermal strains.

**Product:** Abaqus/Standard

### References

---

- “Thermal expansion,” Section 23.1.2 of the Abaqus Analysis User’s Manual
- \*EXPANSION
- “UEXPAN,” Section 4.1.16 of the Abaqus Verification Manual

### Overview

---

User subroutine **UEXPAN**:

- can be used to define incremental thermal strains as functions of temperature, predefined field variables, and state variables;
- is intended for models in which the thermal strains depend on temperature and/or predefined field variables in complex ways or depend on state variables, which can be used and updated in this routine;
- is called at all integration points of elements for which the material or gasket behavior definition contains user-subroutine-defined thermal expansion; and
- is called twice per material point in each iteration during coupled temperature-displacement analysis.

### User subroutine interface

---

```

SUBROUTINE UEXPAN(EXPAN,DEXPANDT,TEMP,TIME,DTIME,PREDEF,
1 DPRED,STATEV,CMNAME,NSTATV,NOEL)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
C
DIMENSION EXPAN(*),DEXPANDT(*),TEMP(2),TIME(2),PREDEF(*),
1 DPRED(*),STATEV(NSTATV)

```

*user coding to define EXPAN, DEXPANDT and update  
STATEV if necessary.*

```

RETURN
END

```

## Variables to be defined

---

### **EXPAN (\*)**

Increments of thermal strain. The number of values to be defined and the order in which they are arranged depend on the type of thermal expansion being defined.

- For isotropic expansion give the isotropic thermal strain increment as the first and only component of the matrix.
- For orthotropic expansion give  $\Delta\epsilon_{11}^{th}$ ,  $\Delta\epsilon_{22}^{th}$ , and  $\Delta\epsilon_{33}^{th}$  as the first, second, and third components of the matrix, respectively.
- For anisotropic expansion give  $\Delta\epsilon_{11}^{th}$ ,  $\Delta\epsilon_{22}^{th}$ ,  $\Delta\epsilon_{33}^{th}$ ,  $\Delta\epsilon_{12}^{th}$ ,  $\Delta\epsilon_{13}^{th}$ , and  $\Delta\epsilon_{23}^{th}$ . Direct components are stored first, followed by shear components in the order presented here. For plane stress only three components of the matrix are needed; give  $\Delta\epsilon_{11}^{th}$ ,  $\Delta\epsilon_{22}^{th}$ , and  $\Delta\epsilon_{12}^{th}$ , as the first, second, and third components, respectively.

### **DEXPANDT (\*)**

Variation of thermal strains with respect to temperature,  $\partial\epsilon^{th}/\partial\theta$ . The number of values and the order in which they are arranged depend on the type of thermal expansion being defined.

- For isotropic expansion give the variation of the isotropic thermal strain with respect to temperature as the first and only component of the matrix.
- For orthotropic expansion give  $\partial\epsilon_{11}^{th}/\partial\theta$ ,  $\partial\epsilon_{22}^{th}/\partial\theta$ , and  $\partial\epsilon_{33}^{th}/\partial\theta$  as the first, second, and third components of the matrix, respectively.
- For anisotropic expansion give  $\partial\epsilon_{11}^{th}/\partial\theta$ ,  $\partial\epsilon_{22}^{th}/\partial\theta$ ,  $\partial\epsilon_{33}^{th}/\partial\theta$ ,  $\partial\epsilon_{12}^{th}/\partial\theta$ ,  $\partial\epsilon_{13}^{th}/\partial\theta$ , and  $\partial\epsilon_{23}^{th}/\partial\theta$ . Direct components are stored first, followed by shear components in the order presented here. For plane stress only three components of the matrix are needed; give  $\partial\epsilon_{11}^{th}/\partial\theta$ ,  $\partial\epsilon_{22}^{th}/\partial\theta$ , and  $\partial\epsilon_{12}^{th}/\partial\theta$ , as the first, second, and third components, respectively.

## Variable that can be updated

---

### **STATEV (NSTATV)**

Array containing the user-defined solution-dependent state variables at this point. Except for coupled temperature-displacement analysis, these are supplied as values at the start of the increment and can be updated to their values at the end of the increment. For coupled temperature-displacement analysis, **UEXPAN** is called twice per material point per iteration. In the first call for a given material point and iteration, the values supplied are those at the start of the increment and can be updated. In the second call for the same material point and iteration, the values supplied are those returned from the first call, and they can be updated again to their values at the end of the increment.

User subroutine **UEXPAN** allows for the incremental thermal strains to be only weakly dependent on the state variables. The Jacobian terms arising from the derivatives of the thermal strains with respect to the state variables are not taken into account.

---

**Variables passed in for information****TEMP (1)**

Current temperature (at the end of the increment).

**TEMP (2)**

Temperature increment.

**TIME (1)**

Step time at the end of the increment.

**TIME (2)**

Total time at the end of the increment.

**DTIME**

Time increment.

**PREDEF (\*)**

Array containing the values of all the user-specified predefined field variables at this point (initial values at the beginning of the analysis and current values during the analysis).

**DPRED (\*)**

Array of increments of predefined field variables.

**CMNAME**

User-specified material name or gasket behavior name, left justified.

**NSTATV**

Number of solution-dependent state variables associated with this material or gasket behavior type (specified when space is allocated for the array; see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NOEL**

User-defined element number.



## 1.1.27 UEXTERNALDB: User subroutine to manage user-defined external databases and calculate model-independent history information.

**Product:** Abaqus/Standard

### Reference

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual

### Overview

---

User subroutine **UEXTERNALDB**:

- is called once each at the beginning of the analysis, at the beginning of each increment, at the end of each increment, and at the end of the analysis (in addition, the subroutine is also called once at the beginning of a restart analysis);
- can be used to communicate between other software and user subroutines within Abaqus/Standard;
- can be used to open external files needed for other user subroutines at the beginning of the analysis and to close those files at the end of the analysis;
- can be used to calculate or read history information at the beginning of each increment. This information can be written to user-defined COMMON block variables or external files for use during the analysis by other user subroutines; and
- can be used to write the current values of the user-calculated history information to external files.

### User subroutine interface

---

```

SUBROUTINE UEXTERNALDB (LOP,LRESTART,TIME,DTIME,KSTEP,KINC)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION TIME(2)
C
user coding to set up the FORTRAN environment, open files, close files,
calculate user-defined model-independent history information,
write history information to external files,
recover history information during restart analyses, etc.
do not include calls to utility routine XIT
C
RETURN
END

```

---

**Variable to be defined**

None.

---

**Variables passed in for information****LOP**

**LOP**=0 indicates that the subroutine is being called at the start of the analysis.

**LOP**=1 indicates that the subroutine is being called at the start of the current analysis increment. The subroutine can be called multiple times at the beginning of an analysis increment if the increment fails to converge and a smaller time increment is required.

**LOP**=2 indicates that the subroutine is being called at the end of the current analysis increment. When **LOP**=2, all information that you need to restart the analysis should be written to external files.

**LOP**=3 indicates that the subroutine is being called at the end of the analysis.

**LOP**=4 indicates that the subroutine is being called at the beginning of a restart analysis. When **LOP**=4, all necessary external files should be opened and properly positioned and all information required for the restart should be read from the external files.

**LRESTART**

**LRESTART**=0 indicates that an analysis restart file is not being written for this increment.

**LRESTART**=1 indicates that an analysis restart file is being written for this increment.

**LRESTART**=2 indicates that an analysis restart file is being written for this increment and that only one increment is being retained per step so that the current increment overwrites the previous increment in the restart file (see “Restarting an analysis,” Section 9.1.1 of the Abaqus Analysis User’s Manual).

**TIME (1)**

Value of current step time.

**TIME (2)**

Value of current total time.

**DTIME**

Time increment.

**KSTEP**

Current step number. When **LOP**=4, **KSTEP** gives the restart step number.

**KINC**

Current increment number. When **LOP**=4, **KINC** gives the restart increment number.

## 1.1.28      **UFIELD:**    User subroutine to specify predefined field variables.

**Product:** Abaqus/Standard

### References

---

- “USDFLD,” Section 1.1.46
- “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual
- \*FIELD
- “**UTEMP, UFIELD, UMASFL, and UPRESS,**” Section 4.1.25 of the Abaqus Verification Manual

### Overview

---

User subroutine **UFIELD**:

- allows you to prescribe predefined field variables at the nodes of a model—the predefined field variables at a node can be updated individually, or a number of field variables at the node can be updated simultaneously;
- is called whenever a user-subroutine-defined field appears;
- ignores any field variable values specified directly;
- can be used to modify field variable values read from a results file; and
- can be used in conjunction with user subroutine **USDFLD** such that the field variables that are passed in from **UFIELD** and interpolated to the material points can be modified (such changes are local to material point values, and nodal field variable values remain unaffected).

### Updating field variables

---

Two different methods are provided for updating field variables.

#### Individual variable updates

By default, only one field variable at a time can be updated in user subroutine **UFIELD**. In this case the user subroutine will be called whenever a current value of a field variable is needed for a node that is listed in the specified field variable definition. This method can be used only for cases in which the field variables are independent of each other.

#### Simultaneous variable updates

For cases in which the field variables depend on each other, multiple (possibly all) field variables at a point can be updated simultaneously in user subroutine **UFIELD**. In this case you must specify the number of field variables to be updated simultaneously at a point, and the user subroutine will be called each time the current field variable values are needed.

**User subroutine interface**

---

```

SUBROUTINE UFIELD (FIELD ,KFIELD ,NSECPT ,KSTEP ,KINC ,TIME ,NODE ,
1 COORDS ,TEMP ,DTEMP ,NFIELD)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION FIELD (NSECPT,NFIELD) , TIME (2) , COORDS (3) ,
1 TEMP (NSECPT) , DTEMP (NSECPT)
C

```

*user coding to define **FIELD***

```

RETURN
END

```

**Variable to be defined**

---

**FIELD (NSECPT ,NFIELD)**

Array of predefined field variable values at node number **NODE**. When updating only one field variable at a time, only the value of the specified field variable (see **KFIELD** below) must be returned. In this case **NFIELD** is passed into user subroutine **UFIELD** with a value of 1, and **FIELD** is thus dimensioned as **FIELD (NSECPT ,1)**. When updating all field variables simultaneously, the values of the specified number of field variables at the point must be returned. In this case **FIELD** is dimensioned as **FIELD (NSECPT ,NFIELD)**, where **NFIELD** is the number of field variables specified and **KFIELD** has no meaning.

If **NODE** is part of any element other than a beam or shell, only one value of each field variable must be returned (**NSECPT**=1). Otherwise, the number of values to be returned depends on the mode of temperature and field variable input selected for the beam or shell section. The following cases are possible:

1. Temperatures and field variables for a beam section are given as values at the points shown in the beam section descriptions. The number of values required, **NSECPT**, is determined by the particular section type specified, as described in “Beam cross-section library,” Section 26.3.9 of the Abaqus Analysis User’s Manual.
2. Temperatures and field variables are given as values at  $n$  equally spaced points through each layer of a shell section. The number of values required, **NSECPT**, is equal to  $n$ .
3. Temperatures and field variables for a beam section are given as values at the origin of the cross-section together with gradients with respect to the 2-direction and, for three-dimensional beams, the 1-direction of the section; or temperatures and field variables for a shell section are given

as values at the reference surface together with gradients through the thickness. The number of values required, **NSECPT**, is 3 for three-dimensional beams, 2 for two-dimensional beams, and 2 for shells. Give the midsurface value first, followed by the first and (if necessary) second gradients, as described in “Beam elements,” Section 26.3 of the Abaqus Analysis User’s Manual, and “Shell elements,” Section 26.6 of the Abaqus Analysis User’s Manual.

Since field variables can also be defined directly, it is important to understand the hierarchy used in situations of conflicting information (see “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual).

When the array **FIELD** is passed into user subroutine **UFIELD**, it will contain either the field variable values from the previous increment or those values obtained from the results file if this method was used. You are then free to modify these values within this subroutine.

## **Variables passed in for information**

---

### **KFIELD**

User-specified field variable number. This variable is meaningful only when updating individual field variables at a time.

### **NFIELD**

User-specified number of field variables to be updated. This variable is meaningful only when updating multiple field variables simultaneously.

### **NSECPT**

Maximum number of section values required for any node in the model. The **NSECPT** can be 2 when only one field variable is specified at some non-beam or non-shell nodes in the model with contact.

### **KSTEP**

Step number.

### **KINC**

Increment number.

### **TIME (1)**

Current value of step time.

### **TIME (2)**

Current total time.

### **NODE**

Node number.

### **COORDS**

An array containing the coordinates of this node. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the node.

## UFIELD

### **TEMP (NSECPT)**

Current temperature at the node. If user subroutines **UTEMP** and **UFIELD** are both used, user subroutine **UTEMP** is processed before user subroutine **UFIELD**.

### **DTEMP (NSECPT)**

Temperature increment at the node.

## 1.1.29 UFLUID: User subroutine to define fluid density and fluid compliance for hydrostatic fluid elements.

**Product:** Abaqus/Standard

### References

---

- “Hydrostatic fluid models,” Section 23.4.1 of the Abaqus Analysis User’s Manual
- \*FLUID PROPERTY
- “UFLUID,” Section 4.1.17 of the Abaqus Verification Manual

### Overview

---

User subroutine **UFLUID**:

- is called for each cavity for which a user-defined fluid constitutive model is being specified;
- is called for every fluid element (“Hydrostatic fluid elements,” Section 29.8.1 of the Abaqus Analysis User’s Manual) and for every fluid link element (“Fluid link elements,” Section 29.8.3 of the Abaqus Analysis User’s Manual) connected to a cavity reference node;
- requires that the fluid density,  $\rho(p, \theta)$ , and the fluid pressure compliance,  $C_p$ , be defined;
- requires that the fluid temperature compliance,  $C_\theta$ , be defined if the routine is to be used in a linear perturbation step and the fluid is subjected to a temperature excursion; and
- ignores any data specified for the fluid constitutive model outside the user subroutine.

### Density and fluid mass

---

At the start of the analysis (prior to the first iteration) the density calculated in user subroutine **UFLUID** (for the initial pressure,  $p_I$ , and temperature,  $\theta_I$ ) is used to calculate the fluid mass from the initial cavity volume. During the analysis the expected cavity volume is calculated from the fluid mass and the density.

### User subroutine interface

---

```

SUBROUTINE UFLUID (RHO, CP, CT, PNEWDT, ENER, PRESS, DPRESS, PRESSI,
1 TEMP, DTEMP, TEMPI, TIME, DTIME, KSTEP, KINC, NONUM, FLNAME, LFLAG)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 FLNAME
DIMENSION TIME(2)

user coding to define RHO, CP, and CT

RETURN
END

```

**Variables to be defined**

---

**RHO**

Fluid density,  $\rho$ , at the end of the increment.

**CP**

Fluid pressure compliance,  $C_p$ , at the end of the increment. For a linear perturbation step this is the base state compliance. Fluid pressure compliance is defined as

$$C_p = \frac{d\rho^{-1}}{dp} = -\rho^{-2} \frac{d\rho}{dp},$$

where  $p$  is the fluid cavity pressure.

**CT**

Fluid temperature compliance,  $C_\theta$ . This variable is needed only if a fluid temperature excursion occurs in a linear perturbation step and is the base state compliance. Fluid temperature compliance is defined as

$$C_\theta = \frac{d\rho^{-1}}{d\theta} = -\rho^{-2} \frac{d\rho}{d\theta},$$

where  $\theta$  is the fluid cavity temperature.

**Variables that can be updated**

---

**PNEWDT**

Ratio of suggested new time increment to the time increment being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **UFLUID**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT**  $\times$  **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT**  $\times$  **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

**ENER**

Energy per unit mass stored in the fluid. This variable is used for energy output only and has no effect on the solution.

---

**Variables passed in for information****PRESS**

Fluid cavity pressure at the end of the increment. For a linear perturbation step this is the base state pressure.

**DPRESS**

Fluid cavity pressure increment. For a linear perturbation step this value is zero.

**PRESSI**

Fluid cavity pressure at the beginning of the analysis.

**TEMP**

Fluid cavity temperature at the end of the increment. For a linear perturbation step this is the base state temperature.

**DTEMP**

Fluid cavity temperature increment. For a linear perturbation step this value is zero.

**TEMPI**

Fluid cavity temperature at the beginning of the analysis.

**TIME (1)**

Current value of step time at the start of the increment.

**TIME (2)**

Current value of total time at the start of the increment.

**DTIME**

Time increment.

**KSTEP**

Step number.

**KINC**

Increment number.

**NONUM**

Cavity reference node number.

**FLNAME**

User-specified fluid property name, left justified.

## **UFLUID**

### **LFLAG**

Linear perturbation flag for the step. If this is a linear perturbation step, **LFLAG**=1. For a general analysis step **LFLAG**=0.

**1.1.30     UFLUIDLEAKOFF: User subroutine to define the fluid leak-off coefficients for pore pressure cohesive elements.**

**Product:** Abaqus/Standard

## References

---

- “Defining the constitutive response of fluid within the cohesive element gap,” Section 29.5.7 of the Abaqus Analysis User’s Manual
- \*FLUID LEAKOFF
- “Propagation of hydraulically driven fracture,” Section 3.3.2 of the Abaqus Verification Manual

## Overview

---

User subroutine **UFLUIDLEAKOFF**:

- can be used to define the fluid leak-off coefficients for pore pressure cohesive elements;
- is called at all material calculation points of elements for which the material definition contains user-defined leak-off coefficients; and
- can include material behavior dependent on field variables or state variables.

## User subroutine interface

---

```

      SUBROUTINE UFLUIDLEAKOFF(PERM,PGRAD,DN,P_INT,P_BOT,P_TOP,
1 ANM,TANG,TIME,DTIME,TEMP,DTEMP,PREDEF,DPRED,C_BOT,C_TOP,
2 DC_BOT,DC_TOP,STATEV,NSTATV,NOEL,NPT,KSTEP,KINC)
C
      INCLUDE 'ABA_PARAM.INC'
C
      CHARACTER*80 CMNAME
      DIMENSION PERM(2),PGRAD(2),ANM(3),TANG(3,2),TIME(2),PREDEF(1),
1 DPRED(1),DC_BOT(3),DC_TOP(3),STATEV(NSTATV)

```

*user coding to define C\_BOT, C\_TOP, DC\_BOT, and DC\_TOP*

```

      RETURN
      END

```

**Variables to be defined**

---

**C\_BOT**

$C_{bot}$ , fluid leak-off coefficient on the bottom side of a pore pressure cohesive element.

**C\_TOP**

$C_{top}$ , fluid leak-off coefficient on the top side of a pore pressure cohesive element.

**DC\_BOT(1)**

$\partial C_{bot} / \partial d$ , where  $d = \text{DN}$ .

**DC\_BOT(2)**

$\partial C_{bot} / \partial p_{int}$ , where  $p_{int} = \text{P\_INT}$ .

**DC\_BOT(3)**

$\partial C_{bot} / \partial p_{bot}$ , where  $p_{bot} = \text{P_BOT}$ .

**DC\_TOP(1)**

$\partial C_{top} / \partial d$ , where  $d = \text{DN}$ .

**DC\_TOP(2)**

$\partial C_{top} / \partial p_{int}$ , where  $p_{int} = \text{P\_INT}$ .

**DC\_TOP(3)**

$\partial C_{top} / \partial p_{top}$ , where  $p_{top} = \text{P_TOP}$ .

**STATEV(NSTATV)**

An array containing the values of the solution-dependent state variables. You define the meaning of these variables. These are passed in as the values at the beginning of the increment and must be returned as the values at the end of the increment. The size of the array is defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

**Variables passed in for information**

---

**PERM(1)**

Fluid permeability.

**PERM(2)**

The derivative of fluid permeability with regard to the opening.

**PGRAD(1)**

The first component of internal pressure gradient.

**PGRAD(2)**

The second component of internal pressure gradient.

**DN**

The relative opening of the element.

**P\_INT**

Internal pressure.

**P\_BOT**

Bottom pressure.

**P\_TOP**

Top pressure.

**ANM**

Normal vector directed from the bottom face toward the top face.

**TANG**

Tangent direction vectors.

**TIME (1)**

Value of step time at the beginning of the current increment.

**TIME (2)**

Value of total time at the beginning of the current increment.

**DTIME**

Time increment.

**TEMP**

Temperature at the start of the increment.

**DTEMP**

Increment of temperature.

**PREDEF**

Array of interpolated values of predefined field variables at this point at the start of the increment, based on the values read in at the nodes.

**DPRED**

Array of increments of predefined field variables.

**NSTATTV**

Number of solution-dependent state variables that are associated with this material type (defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

## **UFLUIDLEAKOFF**

### **NOEL**

Element number.

### **NPT**

Integration point number.

### **KSTEP**

Step number.

### **KINC**

Increment number.

### 1.1.31 UGENS: User subroutine to define the mechanical behavior of a shell section.

**Product:** Abaqus/Standard

#### References

---

- “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual
- \*SHELL GENERAL SECTION

#### Overview

---

User subroutine **UGENS**:

- is used to define the (nonlinear) mechanical behavior of a shell section directly in terms of generalized section quantities;
- requires you to define the section behavior of the shell directly in terms of membrane stresses and forces, curvature changes, and bending moments;
- will be called at all integration points in all shell elements with a general, arbitrary, elastic shell section and a user-subroutine-defined shell section stiffness; and
- can be used with all static or dynamic procedures other than the quasi-static procedure, since that procedure uses automatic time stepping based on the techniques used by Abaqus/Standard to integrate standard creep laws.

#### Storage of membrane and bending components

---

In the force and strain arrays and in the matrix **DDNDDE**, direct membrane terms are stored first, followed by the shear membrane term, and then the direct and shear bending terms. Only active components are stored, so the number of entries depends on the element type (see Table 1.1.31–1).

**Table 1.1.31–1** Active section force/moment components.

Element type	Force and moment components
Three-dimensional shells (S4R, S8R, S8R5, etc.) and axisymmetric shells with asymmetric deformation (SAXA1N, SAXA2N)	$N_{11}, N_{22}, N_{12}, M_{11}, M_{22}, M_{12}$
Axisymmetric shells (SAX1, SAX2, etc)	$N_{11}, N_{22}, M_{11}, M_{22}$

There are **NDI** direct membrane and **NSHR** shear membrane components and **NDI** direct bending and **NSHR** shear bending components: a total of **NSECV** components. The order of the components is defined in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual.

Engineering measures of shear membrane strain ( $\gamma_{12}$ ) and twist ( $K_{12}$ ) are used.

## **Increments for which only the section stiffness can be defined**

---

Abaqus/Standard passes zero strain increments into user subroutine **UGENS** to start the first increment of all the steps and all increments of steps for which you have suppressed extrapolation in time from the previous incremental solution (“Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual). In this case you can define only the section stiffness (**DDNDDE**).

## **Stability**

---

You should ensure that the integration scheme coded in this routine is stable—no direct provision is made to include a stability limit in the time stepping scheme based on the calculations in **UGENS**.

## **Convergence rate**

---

**DDNDDE** must be defined accurately if rapid convergence of the overall Newton scheme is to be achieved. In most cases the accuracy of this definition is the most important factor governing the convergence rate. Unsymmetric equation solution is as much as four times as expensive as the corresponding symmetric system. Therefore, if the section stiffness matrix (**DDNDDE**) is only slightly unsymmetric, it may be computationally less expensive to use a symmetric approximation and accept a slightly slower rate of convergence.

## **Use with shells that have transverse shear and/or hourglass stiffness**

---

If user subroutine **UGENS** is used to describe the section behavior of shells with transverse shear, you must define the transverse shear stiffness (see “Defining the transverse shear stiffness” in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual).

If user subroutine **UGENS** is used to describe the section behavior of shells with hourglass stiffness, you must define the hourglass stiffness parameter for hourglass control based on total stiffness (see “Specifying nondefault hourglass control parameters for reduced-integration shell elements” in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual). The hourglass stiffness parameter is not required for enhanced hourglass control, but you can define a scaling factor for the stiffness associated with the drill degree of freedom (rotation about the surface normal).

## **Use with continuum shell elements**

---

User subroutine **UGENS** cannot be used to describe the section behavior of continuum shell elements.

## **User subroutine interface**

---

```
SUBROUTINE UGENS (DDNDDE , FORCE , STATEV , SSE , SPD , PNEWDT , STRAN ,
1 DSTRAN , TSS , TIME , DTIME , TEMP , DTEMP , PREDEF , DPRED , CENAME , NDI ,
2 NSHR , NSECV , NSTATV , PROPS , JPROPS , NPROPS , NJPROP , COORDS , CELENT ,
3 THICK , DFGRD , CURV , BASIS , NOEL , NPT , KSTEP , KINC , NIT , LINPER)
```

```

C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CENAME
DIMENSION DDNDDE (NSECV,NSECV) ,FORCE (NSECV) ,STATEV (NSTATV) ,
1 STRAN (NSECV) ,DSTRAN (NSECV) ,TSS (2) ,TIME (2) ,PREDEF (*),
2 DPRED (*), PROPS (*), JPROPS (*), COORDS (3) ,DFGRD (3,3) ,
3 CURV (2,2) ,BASIS (3,3)

user coding to define DDNDDE, FORCE, STATEV, SSE, PNEWDT

RETURN
END

```

## **Variables to be defined**

---

### **DDNDDE (NSECV, NSECV)**

Section stiffness matrix of the shell section,  $\partial\mathbf{N}/\partial\mathbf{E}$ , where  $\mathbf{N}$  are the section forces and moments on the shell section and  $\mathbf{E}$  are the generalized section strains in the shell. **DDNDDE (I, J)** defines the change in the **I**th force component at the end of the time increment caused by an infinitesimal perturbation of the **J**th component of the section strain increment array. The size of this matrix depends on the values of **NSECV** (see below for details).

Unless you invoke the unsymmetric equation solution capability in the general shell section definition (“Defining whether or not the section stiffness matrices are symmetric” in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual), Abaqus/Standard will use only the symmetric part of **DDNDDE**. The symmetric part of the matrix is calculated by taking one half the sum of the matrix and its transpose.

### **FORCE (NSECV)**

This array is passed in as the forces and moments per unit length on the shell surface at the beginning of the increment and must be updated in this routine to be the forces and moments at the end of the increment.

### **STATEV (NSTATV)**

An array containing the solution-dependent state variables. These are passed in as the values at the beginning of the increment and must be returned as the values at the end of the increment.

### **SSE, SPD**

Elastic strain energy and plastic dissipation, respectively. These are passed in as the values at the beginning of the increment and should be updated to the corresponding energy values at the end of the increment. These values have no effect on the solution; they are used for the energy output.

## Variable that can be updated

---

### PNEWDT

Ratio of suggested new time increment to the time increment being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **UGENS**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

## Variables passed in for information

---

### STRAN (NSECV)

An array containing the generalized section strains (membrane strains and curvature changes) at the beginning of the increment. The size of this array depends on the value of **NSECV** (see below for details).

### DSTRAN (NSECV)

Array of generalized section strain increments.

### TSS (2)

Array containing the transverse shear strains.

### TIME (1)

Value of step time at the beginning of the current increment.

### TIME (2)

Value of total time at the beginning of the current increment.

### DTIME

Time increment.

### TEMP

Temperature at the start of the increment.

**DTEMP**

Increment of temperature.

**PREDEF**

Array of interpolated values of predefined field variables at this point at the start of the increment, based on the values read in at the nodes.

**DPRED**

Array of increments of predefined field variables.

**CENAME**

User-specified element set name associated with this section, left justified.

**NDI**

Number of direct force components at this point.

**NSHR**

Number of shear force components at this point.

**NSECV**

Size of the force and strain component arrays.

**NSTATV**

User-defined number of solution-dependent state variables associated with this section (“Defining the number of solution-dependent variables that must be stored for the section” in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual).

**PROPS (NPROPS)**

A floating point array containing the **NPROPS** real property values defined for use with this section.

**JPROPS (NJPROP)**

An integer array containing the **NJPROP** integer property values defined for use with this section.

**NPROPS**

User-defined number of real property values associated with this section (“Defining the section properties” in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual).

**NJPROP**

User-defined number of integer property values associated with the element (“Defining the section properties” in “Using a general shell section to define the section behavior,” Section 26.6.6 of the Abaqus Analysis User’s Manual).

**COORDS**

An array containing the current coordinates of this integration point.

**CELENT**

Characteristic element length in the reference surface.

**THICK**

Original section thickness.

**DFGRD (3 , 3)**

An array containing the components of the midsurface deformation gradient,  $\bar{f}_{ij}$ . The deformation gradient curvature tensor is available for finite-strain shells (S3/S3R, S4, S4R, SAXs, and SAXAs); it is not available for small-strain shells.

The deformation gradient is stored as a  $3 \times 3$  matrix with component equivalence **DFGRD (I , J)  $\Leftrightarrow$   $\bar{f}_{ij}$** .  $\bar{f}_{\alpha\beta}$  (Greek subscripts range from 1 to 2) are the in-plane components of the deformation gradient, and  $\bar{f}_{33}$  is the thickness change component. The components,  $\bar{f}_{\alpha 3}$ , are the transverse shear strains scaled by  $\bar{f}_{33}$ . The remaining components,  $\bar{f}_{3\beta}$ , are all zero.

The tensor is provided in the local shell coordinate system.

**CURV (2 , 2)**

An array containing the midsurface curvature tensor,  $b_{\alpha\beta}$ . The curvature tensor is available for finite-strain shells (S3/S3R, S4, S4R, SAXs, and SAXAs); it is not available for small-strain shells.

The curvature tensor is stored as a  $2 \times 2$  matrix with component equivalence **CURV (I , J)  $\Leftrightarrow$   $b_{\alpha\beta}$** .

The tensor is provided in the local shell coordinate system.

**BASIS (3 , 3)**

An array containing the direction cosines of the shell local surface coordinate system. **BASIS (1 , 1)**, **BASIS (2 , 1)**, and **BASIS (3 , 1)** give the (1, 2, 3) components of the first local direction, etc. The first two directions are in the plane of the element surface, and the third direction is the normal. The conventions for local directions on shell surfaces are defined in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual. You can redefine the local system; see “Orientations,” Section 2.2.5 of the Abaqus Analysis User’s Manual.

**NOEL**

Element number.

**NPT**

Integration point number.

**KSTEP**

Step number.

**KINC**

Increment number.

**NIT**

Iteration number. **NIT=0** during the first assembly of the system matrix in any increment.

**LINPER**

Linear perturbation flag. **LINPER**=1 if the step is a linear perturbation step. **LINPER**=0 if the step is a general step.



**1.1.32      UHARD: User subroutine to define the yield surface size and hardening parameters for isotropic plasticity or combined hardening models.**

**Product:** Abaqus/Standard

## References

---

- “Classical metal plasticity,” Section 20.2.1 of the Abaqus Analysis User’s Manual
- “Models for metals subjected to cyclic loading,” Section 20.2.2 of the Abaqus Analysis User’s Manual
- \*CYCLIC HARDENING
- \*PLASTIC

## Overview

---

User subroutine **UHARD**:

- is called at all material calculation points of elements for which the material definition includes user-defined isotropic hardening or cyclic hardening for metal plasticity;
- can be used to define a material’s isotropic yield behavior;
- can be used to define the size of the yield surface in a combined hardening model;
- can include material behavior dependent on field variables or state variables; and
- requires, when appropriate, that the values of the derivatives of the yield stress (or yield surface size in combined hardening models) be defined with respect to the strain, strain rate, and temperature.

## User subroutine interface

---

```

SUBROUTINE UHARD (SYIELD , HARD , EQPLAS , EQPLASRT , TIME , DTIME , TEMP ,
1      DTEMP , NOEL , NPT , LAYER , KSPT , KSTEP , KINC , CMNAME , NSTATV ,
2      STATEV , NUMFIELDV , PREDEF , DPRED , NUMPROPS , PROPS )
C
C      INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
DIMENSION HARD (3) , STATEV(NSTATV) , TIME (*) ,
$           PREDEF (NUMFIELDV) , DPRED (*) , PROPS (*)

user coding to define SYIELD , HARD (1) , HARD (2) , HARD (3)

RETURN
END

```

**Variables to be defined**

---

**SYIELD**

$\sigma^0$ . Yield stress for isotropic plasticity. Yield surface size for combined hardening.

**HARD (1)**

Variation of **SYIELD** with respect to the equivalent plastic strain,  $\partial\sigma^0/\partial\bar{\varepsilon}^{pl}$ .

**HARD (2)**

Variation of **SYIELD** with respect to the equivalent plastic strain rate,  $\partial\sigma^0/\partial\dot{\bar{\varepsilon}}^{pl}$ . This quantity is not used with the combined hardening model.

**HARD (3)**

Variation of **SYIELD** with respect to temperature,  $\partial\sigma^0/\partial\theta$ . This quantity is required only in adiabatic and fully coupled temperature-displacement analyses.

**STATEV (NSTATV)**

Array containing the user-defined solution-dependent state variables at this point. These are supplied as values at the beginning of the increment or as values updated by other user subroutines (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual) and must be returned as values at the end of the increment.

**Variables passed in for information**

---

**EQPLAS**

Equivalent plastic strain,  $\bar{\varepsilon}^{pl}$ .

**EQPLASRT**

Equivalent plastic strain rate,  $\dot{\bar{\varepsilon}}^{pl}$ .

**TIME (1)**

Value of step time at the beginning of the current increment.

**TIME (2)**

Value of total time at the beginning of the current increment.

**DTIME**

Time increment.

**TEMP**

Temperature at the beginning of the increment.

**DTEMP**

Increment of temperature.

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**KSTEP**

Step number.

**KINC**

Increment number.

**CMNAME**

User-specified material name, left justified.

**NSTATTV**

User-specified number of solution-dependent state variables associated with this material (“Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NUMFIELDV**

Number of field variables.

**PREDEF (NUMFIELDV)**

Array of interpolated values of predefined field variables at this material point at the start of the increment based on the values read in at the nodes (initial values at the beginning of the analysis and current values during the analysis).

**DPRED (NUMFIELDV)**

Array of increments of predefined field variables at this material point for this increment; this includes any values updated by user subroutine **USDFLD**.

**NPROPS**

Number of hardening properties entered for this user-defined hardening definition.

**PROPS (NPROPS)**

Array of hardening properties entered for this user-defined hardening definition.



### 1.1.33 UHYPEL: User subroutine to define a hypoelastic stress-strain relation.

**Product:** Abaqus/Standard

#### References

---

- “Hypoelastic behavior,” Section 19.4.1 of the Abaqus Analysis User’s Manual
- \*HYPOELASTIC

#### Overview

---

User subroutine **UHYPEL**:

- can be used to define isotropic hypoelastic material behavior, thus requiring the definition of Young’s modulus,  $E$ , and Poisson’s ratio,  $\nu$ ;
- is called at all material calculation points of elements for which the material definition contains user-defined hypoelastic behavior;
- can be used in conjunction with user subroutine **USDFLD** to redefine any field variables that are passed in (see “USDFLD,” Section 1.1.46); and
- ignores any data specified outside the user subroutine for the associated hypoelastic material definition.

#### Special considerations for various element types

---

There are several special considerations that need to be noted.

##### Beams and shells that calculate transverse shear energy

When **UHYPEL** is used to define the material response of shell or beam elements that calculate transverse shear energy, Abaqus/Standard cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element’s transverse shear stiffness. See “Shell section behavior,” Section 26.6.4 of the Abaqus Analysis User’s Manual, and “Choosing a beam element,” Section 26.3.3 of the Abaqus Analysis User’s Manual, for guidelines on choosing this stiffness.

##### Elements with hourgassing modes

If this capability is used to describe the material of elements with hourgassing modes, you must define the hourglass stiffness for hourglass control based on the total stiffness approach. The hourglass stiffness is not required for enhanced hourglass control, but you can define a scaling factor for the stiffness associated with the drill degree of freedom (rotation about the surface normal). See “Section controls,” Section 24.1.4 of the Abaqus Analysis User’s Manual.

**User subroutine interface**

---

```

SUBROUTINE UHYPEL (E,GNU,STRAIN,NDI,NSHR,EINV1,EINV2,EINV3,
1 COORDS,NOEL,TEMP,PREDEF,CMNAME)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
C
DIMENSION STRAIN(*),COORDS(3),PREDEF(*)

user coding to define E and GNU

RETURN
END

```

**Variables to be defined**

---

**E**

Young's modulus.

**GNU**

Poisson's ratio.

**Variables passed in for information**

---

**STRAIN**

Array containing the total (elastic) strains, ( $\varepsilon$ ).

**NDI**

Number of direct strain components at this point.

**NSHR**

Number of shear strain components at this point.

**EINV1**

$I_1 = \text{trace}(\varepsilon)$ , the first strain invariant.

**EINV2**

$I_2 = 1/2(\varepsilon : \varepsilon - I_1^2)$ , the second strain invariant.

**EINV3**

$I_3 = \det(\varepsilon)$ , the third strain invariant.

**COORDS**

An array containing the coordinates of the material point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**NOEL**

Element number.

**TEMP**

Current temperature at this point.

**PREDEF**

An array containing current values of the predefined field variables at this point (initial values at the beginning of the analysis and current values during the analysis).

**CMNAME**

User-specified material name, left justified.



## 1.1.34 UHYPER: User subroutine to define a hyperelastic material.

**Product:** Abaqus/Standard

### References

---

- “Hyperelastic behavior of rubberlike materials,” Section 19.5.1 of the Abaqus Analysis User’s Manual
- \*HYPERELASTIC
- “UMAT and UHYPER,” Section 4.1.21 of the Abaqus Verification Manual

### Overview

---

User subroutine **UHYPER**:

- can be used to define the strain energy potential for isotropic hyperelastic material behavior;
- is called at all material calculation points of elements for which the material definition contains user-defined hyperelastic behavior;
- can include material behavior dependent on field variables or state variables; and
- requires that the values of the derivatives of the strain energy density function of the hyperelastic material be defined with respect to the strain invariants.

### Special considerations for various element types

---

There are several special considerations that need to be noted.

#### Shells that calculate transverse shear energy

When **UHYPER** is used to define the material response of shell elements that calculate transverse shear energy, Abaqus/Standard cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element’s transverse shear stiffness. See “Shell section behavior,” Section 26.6.4 of the Abaqus Analysis User’s Manual, for guidelines on choosing this stiffness.

#### Elements with hourgassing modes

If this capability is used to describe the material of elements with hourgassing modes, you must define the hourglass stiffness for hourglass control based on the total stiffness approach. The hourglass stiffness is not required for enhanced hourglass control, but you can define a scaling factor for the stiffness associated with the drill degree of freedom (rotation about the surface normal). See “Section controls,” Section 24.1.4 of the Abaqus Analysis User’s Manual.

### User subroutine interface

---

```
SUBROUTINE UHYPER(BI1,BI2,AJ,U,UI1,UI2,UI3,TEMP,NOEL,
1 CMNAME,INCMPFLAG,NUMSTATEV,STATEV,NUMFIELDV,FIELDV,
```

```

2 FIELDVINC ,NUMPROPS ,PROPS)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
DIMENSION U(2) ,UI1(3) ,UI2(6) ,UI3(6) ,STATEV(*) ,FIELDV(*) ,
2 FIELDVINC(*) ,PROPS(*)

```

*user coding to define U,UI1,UI2,UI3,STATEV*

```

RETURN
END

```

## Variables to be defined

---

### U(1)

$U$ , strain energy density function. For a compressible material, at least one derivative involving  $J$  should be nonzero. For an incompressible material, all derivatives involving  $J$  will be ignored. The strain invariants— $\bar{I}_1$ ,  $\bar{I}_2$ , and  $J$ —are defined in “Hyperelastic behavior of rubberlike materials,” Section 19.5.1 of the Abaqus Analysis User’s Manual.

### U(2)

$\tilde{U}_{dev}$ , the deviatoric part of the strain energy density of the primary material response. This quantity is needed only if the current material definition also includes Mullins effect (see “Mullins effect,” Section 19.6.1 of the Abaqus Analysis User’s Manual).

### UI1(1)

$$\partial U / \partial \bar{I}_1.$$

### UI1(2)

$$\partial U / \partial \bar{I}_2.$$

### UI1(3)

$$\partial U / \partial J.$$

### UI2(1)

$$\partial^2 U / \partial \bar{I}_1^2.$$

### UI2(2)

$$\partial^2 U / \partial \bar{I}_2^2.$$

### UI2(3)

$$\partial^2 U / \partial J^2.$$

**UI2 (4)**  
 $\partial^2 U / \partial \bar{I}_1 \partial \bar{I}_2.$

**UI2 (5)**  
 $\partial^2 U / \partial \bar{I}_1 \partial J.$

**UI2 (6)**  
 $\partial^2 U / \partial \bar{I}_2 \partial J.$

**UI3 (1)**  
 $\partial^3 U / \partial \bar{I}_1^2 \partial J.$

**UI3 (2)**  
 $\partial^3 U / \partial \bar{I}_2^2 \partial J.$

**UI3 (3)**  
 $\partial^3 U / \partial \bar{I}_1 \partial \bar{I}_2 \partial J.$

**UI3 (4)**  
 $\partial^3 U / \partial \bar{I}_1 \partial J^2.$

**UI3 (5)**  
 $\partial^3 U / \partial \bar{I}_2 \partial J^2.$

**UI3 (6)**  
 $\partial^3 U / \partial J^3.$

#### STATEEV

Array containing the user-defined solution-dependent state variables at this point. These are supplied as values at the start of the increment or as values updated by other user subroutines (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual) and must be returned as values at the end of the increment.

---

#### Variables passed in for information

**BI1**  
 $\bar{I}_1.$

**BI2**  
 $\bar{I}_2.$

**AJ**  
 $J.$

**TEMP**  
 Current temperature at this point.

**NOEL**

Element number.

**CMNAME**

User-specified material name, left justified.

**INCMPFLAG**

Incompressibility flag defined to be 1 if the material is specified as incompressible or 0 if the material is specified as compressible.

**NUMSTATEV**

User-defined number of solution-dependent state variables associated with this material (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NUMFIELDV**

Number of field variables.

**FIELDV**

Array of interpolated values of predefined field variables at this material point at the end of the increment based on the values read in at the nodes (initial values at the beginning of the analysis and current values during the analysis).

**FIELDVINC**

Array of increments of predefined field variables at this material point for this increment; this includes any values updated by the user subroutine **USDFLD**.

**NUMPROPS**

Number of material properties entered for this user-defined hyperelastic material.

**PROPS**

Array of material properties entered for this user-defined hyperelastic material.

### 1.1.35 UINTER: User subroutine to define surface interaction behavior for contact surfaces.

**Product:** Abaqus/Standard

#### References

---

- “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual
- \*SURFACE INTERACTION
- “**UINTER**,” Section 4.1.20 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UINTER**:

- is called at points on the slave surface of a contact pair with a user-defined constitutive model defining the interaction between the surfaces;
- can be used to define the mechanical (normal and shear) and thermal (heat flux) interactions between surfaces;
- can be used when the normal surface behavior (contact pressure versus overclosure) models (“Contact pressure-overclosure relationships,” Section 33.1.2 of the Abaqus Analysis User’s Manual) or the extended versions of the classical Coulomb friction model (“Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual) are too restrictive and a more complex definition of normal and shear transmission between contacting surfaces, including damping properties, are required;
- must provide the entire definition of the mechanical and the thermal interaction between the contacting surfaces (hence, no additional surface behaviors can be specified in conjunction with this capability);
- can provide the entire definition of viscous and structural damping for interaction between the contacting surfaces for direct steady-state dynamic analysis;
- can use and update solution-dependent state variables; and
- is not available for contact elements.

#### User subroutine interface

---

```
SUBROUTINE UINTER(STRESS, DDSDDR, DVISCOUS, DSTRUCTURAL, FLUX, DDFDDT,
 1 DDSDDT, DDFDDR, STATEV, SED, SFD, SPD, SVD, SCD, PNEWDT, RDISP,
 2 DRDISP,
 3 TEMP, DTEMP, PREDEF, DPRED, TIME, DTIME, FREQR, CINAME, SLNAME,
 4 MSNAME,
```

```

5 PROPS , COORDS , ALOCALDIR , DROT , AREA , CHRLNGTH , NODE , NDIR , NSTATV ,
6 NPRED , NPROPS , MCRD , KSTEP , KINC , KIT , LINPER , LOPENCLOSE , LSTATE ,
7 LSDI , LPRINT)

C
INCLUDE 'ABA_PARAM.INC'

C
CHARACTER*80 CINAME , SLNAME , MSNAME
DIMENSION STRESS (NDIR) , DDSDDR (NDIR,NDIR) , FLUX (2) , DDFDDT (2,2) ,
1 DDSDDT (NDIR,2) , DDFDDR (2,NDIR) , STATEV (NSTATV) ,
2 RDISP (NDIR) , DRDISP (NDIR) , TEMP (2) , DTEMP (2) , PREDEF (2,NPRED) ,
3 DPRED (2,NPRED) , TIME (2) , PROPS (NPROPS) , COORDS (MCRD) ,
4 ALOCALDIR (3,3) , DROT (2,2) , DVISCOUS (NDIR,NDIR) ,
5 DSTRUCTURAL (NDIR,NDIR)

user coding to define STRESS , DDSDDR, FLUX, DDFDDT,
DDSDDT, DDFDDR,
and, optionally, STATEV, SED, SFD, SPD, SVD, SCD, PNEWDT,
LOPENCLOSE, LSTATE, LSDI, DVISCOUS, DSTRUCTURAL
```

**RETURN****END**

## **Variables to be defined**

---

### **STRESS (NDIR)**

This array is passed in as the stress between the slave and master surfaces at the beginning of the increment and must be updated in this routine to be the stress at the end of the increment. The stress must be defined in a local coordinate system (see **ALOCALDIR**). This variable must be defined for a stress/displacement or a fully coupled temperature-displacement analysis. The sign convention for stresses is that a positive stress indicates compression across contact surfaces, while a negative stress indicates tension.

### **DDSDDR (NDIR, NDIR)**

Interface stiffness matrix. **DDSDDR (I, J)** defines the change in the Ith stress component at the end of the time increment caused by an infinitesimal perturbation of the Jth component of the relative displacement increment array. Unless you invoke the unsymmetric equation solution capability in the contact property model definition (“Use with the unsymmetric equation solver in Abaqus/Standard” in “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual), Abaqus/Standard will use only the symmetric part of **DDSDDR**. For a particular off-diagonal (I,J) entry, the symmetrization is done by halving the sum of (I,J) and (J,I) components. **DDSDDR** must be defined for a stress/displacement or a fully coupled temperature-displacement analysis to ensure proper convergence characteristics.

**FLUX (2)**

Magnitude of the heat flux flowing into the slave and master surfaces, respectively. This array is passed in as the value at the beginning of the increment and must be updated to the flux at the end of the increment. The convention for defining the flux is that a positive flux indicates heat flowing into a surface, while a negative flux indicates heat flowing out of the surface. This variable must be defined for a heat transfer or a fully coupled temperature-displacement analysis. The sum of these two flux terms represents the heat generated in the interface, and the difference in these flux terms represents the heat conducted through the interface.

**DDFDDT (2, 2)**

The negative of the variation of the flux at the two surfaces with respect to their respective temperatures, for a fixed relative displacement. This variable must be defined for a heat transfer or a fully coupled temperature-displacement analysis to ensure proper convergence characteristics. The entries in the first row contain the negatives of the derivatives of **FLUX (1)** with respect to **TEMP (1)** and **TEMP (2)**, respectively. The entries in the second row contain the negatives of the corresponding derivatives of **FLUX (2)**.

**DDSDDT (NDIR, 2)**

Variation of the stress with respect to the temperatures of the two surfaces for a fixed relative displacement. This variable is required only for thermally coupled elements (in a fully coupled temperature-displacement analysis), in which the stress is a function of the surface temperatures. **DDSDDT (NDIR, 1)** corresponds to the slave surface, and **DDSDDT (NDIR, 2)** corresponds to the master surface.

**DDFDDR (2, NDIR)**

Variation of the flux with respect to the relative displacement between the two surfaces. This variable is required only for thermally coupled elements (in a fully coupled temperature-displacement analysis), in which the flux is a function of the relative displacement. **DDFDDR (1, NDIR)** corresponds to the slave surface, and **DDFDDR (2, NDIR)** corresponds to the master surface.

---

**Variables that can be updated****DVISCous (NDIR, NDIR)**

Interface viscous damping matrix that can be used only in direct steady-state dynamic analysis. **DVISCous (I, J)** defines an element in the material viscous damping matrix at the current frequency. Abaqus/Standard requires that this element is defined as a damping value for each of the (I, J) entries times the current frequency value **FREQR** obtained from the argument list.

Unless you invoke the unsymmetric equation solution capability in the contact property model definition (“Use with the unsymmetric equation solver in Abaqus/Standard” in “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual), Abaqus/Standard uses only the symmetric part of **DVISCous**. For a particular off-diagonal (I, J) entry the symmetrization is done by halving the sum of the (I, J) and (J, I) components.

**DSTRUCTURAL (NDIR, NDIR)**

Interface structural damping matrix that can be used only in direct steady-state dynamic analysis.

**DSTRUCTURAL (I, J)** defines an element in the material structural damping matrix.

Unless you invoke the unsymmetric equation solution capability in the contact property model definition (“Use with the unsymmetric equation solver in Abaqus/Standard” in “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual), Abaqus/Standard uses only the symmetric part of **DSTRUCTURAL**. For a particular off-diagonal (I, J) entry the symmetrization is done by halving the sum of the (I, J) and (J, I) components.

**STATEV (NSTATV)**

An array containing the solution-dependent state variables. These are passed in as values at the beginning of the increment and must be returned as values at the end of the increment. You define the number of available state variables as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

**SED**

This variable is passed in as the value of the elastic energy density at the start of the increment and should be updated to the elastic energy density at the end of the increment. This variable is used for output only and has no effect on other solution variables. It contributes to the output variable ALLSE.

**SFD**

This variable should be defined as the incremental frictional dissipation. The units are energy per unit area. This variable is used for output only and has no effect on other solution variables. It contributes to the output variables ALLFD and SFDR (and related variables). For computing its contribution to SFDR, **SFD** is divided by the time increment.

**SPD**

This variable should be defined as the incremental dissipation due to plasticity effects in the interfacial constitutive behavior. The units are energy per unit area. This variable is used for output only and has no effect on other solution variables. It contributes to the output variable ALLPD.

**SVD**

This variable should be defined as the incremental dissipation due to viscous effects in the interfacial constitutive behavior. The units are energy per unit area. This variable is used for output only and has no effect on other solution variables. It contributes to the output variable ALLVD.

**SCD**

This variable should be defined as the incremental dissipation due to creep effects in the interfacial constitutive behavior. The units are energy per unit area. This variable is used for output only and has no effect on other solution variables. It contributes to the output variable ALLCD.

**PNEWDT**

Ratio of suggested new time increment to the time increment currently being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **UINTER**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** greater than 1.0 will be ignored and values of **PNEWDT** less than 1.0 will cause the job to terminate.

**LOPENCLOSE**

An integer flag that is used to track the contact status in situations where user subroutine **UINTER** is used to model standard contact between two surfaces, like the default hard contact model in Abaqus/Standard. It comes in as the value at the beginning of the current iteration and should be set to the value at the end of the current iteration. It is set to -1 at the beginning of the analysis before **UINTER** is called. You should set it to 0 to indicate an open status and to 1 to indicate a closed status. A change in this flag from one iteration to the next will have two effects. It will result in output related to a change in contact status if you request a detailed contact printout in the message file (“The Abaqus/Standard message file” in “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). In addition, it will also trigger a severe discontinuity iteration. Any time this flag is reset to a value of -1, Abaqus/Standard assumes that the flag is not being used. A change in this flag from -1 to another value or vice versa will not have any of the above effects.

**LSTATE**

An integer flag that should be used in non-standard contact situations where a simple open/close status is not appropriate or enough to describe the state. It comes in as the value at the beginning of the current iteration and should be set to the value at the end of the current iteration. It is set to -1 at the beginning of the analysis before **UINTER** is called. It can be assigned any user-defined integer value, each corresponding to a different state. You can track changes in the value of this flag and use it to output appropriate diagnostic messages to the message file (unit 7). You may choose to output diagnostic messages only when a detailed contact printout is requested (“The Abaqus/Standard message file” in “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). In the latter case, the **LPRINT** parameter is useful. In conjunction with the **LSTATE** flag, you may also utilize the **LSDI** flag to trigger a severe discontinuity iteration any time the state changes from one iteration to the next. Any time this flag is reset to a value of -1, Abaqus/Standard assumes that the flag is not being used.

**LSDI**

This flag is set to 0 before each call to **UINTER** and should be set to 1 if the current iteration should be treated as a severe discontinuity iteration. This would typically be done in non-standard contact situations based on a change in the value of the **LSTATE** flag from one iteration to the next. The use of this flag has no effect when the **LOPENCLOSE** flag is also used. In that case, severe discontinuity iterations are determined based on changes in the value of **LOPENCLOSE** alone.

---

**Variables passed in for information****RDISP (NDIR)**

An array containing the current relative positions between the two surfaces at the end of the increment. The first component is the relative position of the point on the slave surface, with respect to the master surface, in the normal direction. The second and third components, if applicable, are the accumulated incremental relative tangential displacements, measured from the beginning of the analysis. For the relative position in the normal direction a negative quantity represents an open status, while a positive quantity indicates penetration into the master surface. For open points on the slave surface for which no pairing master is found, the first component is a very large negative number ( $-1 \times 10^{36}$ ). The local directions in which the relative displacements are defined are stored in **ALOCALDIR**.

**DRDISP (NDIR)**

An array containing the increments in relative positions between the two surfaces.

**TEMP (2)**

Temperature at the end of the increment at a point on the slave surface and the opposing master surface, respectively.

**DTEMP (2)**

Increment in temperature at the point on the slave surface and the opposing master surface, respectively.

**PREDEF (2 , NPRED)**

An array containing pairs of values of all the predefined field variables at the end of the current increment (initial values at the beginning of the analysis and current values during the analysis). The first value in a pair, **PREDEF (1 , NPRED)**, corresponds to the value at the point on the slave surface, and the second value, **PFREDEF (2 , NPRED)**, corresponds to the value of the field variable at the nearest point on the opposing surface.

**DPRED (2 , NPRED)**

Array of increments in predefined field variables.

**TIME (1)**

Value of step time at the end of the increment.

**TIME (2)**

Value of total time at the end of the increment.

**DTIME**

Current increment in time.

**FREQR**

Current frequency for direct steady-state dynamic analysis in rad/time.

**CINAME**

User-specified surface interaction name, left justified.

**SLNAME**

Slave surface name.

**MSNAME**

Master surface name.

**PROPS (NPROPS)**

User-specified array of property values to define the interfacial constitutive behavior between the contacting surfaces.

**COORDS (MCRD)**

An array containing the current coordinates of this point.

**ALOCALDIR (3 , 3)**

An array containing the direction cosines of the local surface coordinate system. The directions are stored in columns. For example, **ALOCALDIR (1 , 1)**, **ALOCALDIR (2 , 1)**, and **ALOCALDIR (3 , 1)** give the (1, 2, 3) components of the normal direction. Thus, the first direction is the normal direction to the surface, and the remaining two directions are the slip directions in the plane of the surface. The local system is defined by the geometry of the master surface. The convention for the local directions is the same as the convention in situations where the model uses the built-in contact capabilities in Abaqus/Standard (described in “Contact formulations in Abaqus/Standard,” Section 34.1.1 of the Abaqus Analysis User’s Manual, for the tangential directions).

**DROT (2 , 2)**

Rotation increment matrix. For contact with a three-dimensional rigid surface, this matrix represents the incremental rotation of the surface directions relative to the rigid surface. It is provided so that vector- or tensor-valued state variables can be rotated appropriately in this subroutine. Relative displacement components are already rotated by this amount before **UINTER** is called. This matrix is passed in as a unit matrix for two-dimensional and axisymmetric contact problems.

**AREA**

Surface area associated with the contact point.

**CHRLNGTH**

Characteristic contact surface face dimension.

## **UINTER**

### **NODE**

User-defined global slave node number (or internal node number for models defined in terms of an assembly of part instances) involved with this contact point. Corresponds to the predominant slave node of the constraint if the surface-to-surface contact formulation is used.

### **NDIR**

Number of force components at this point.

### **NSTATV**

Number of solution-dependent state variables.

### **NPRED**

Number of predefined field variables.

### **NPROPS**

User-defined number of property values associated with this interfacial constitutive model (“Interfacial constants” in “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual).

### **MCRD**

Number of coordinate directions at the contact point.

### **KSTEP**

Step number.

### **KINC**

Increment number.

### **KIT**

Iteration number. **KIT**=0 for the first assembly, **KIT**=1 for the first recovery/second assembly, **KIT**=2 for the second recovery/third assembly, and so on.

### **LINPER**

Linear perturbation flag. **LINPER**=1 if the step is a linear perturbation step. **LINPER**=0 if the step is a general step. For a linear perturbation step, the inputs to user subroutine **UINTER** represent perturbation quantities about the base state. The user-defined quantities in **UINTER** are also perturbation quantities. The Jacobian terms should be based on the base state. No change in contact status should occur during a linear perturbation step.

### **LPRINT**

This flag is equal to 1 if a detailed contact printout to the message file is requested and 0 otherwise (“The Abaqus/Standard message file” in “Output,” Section 4.1.1 of the Abaqus Analysis User’s Manual). This flag can be used to print out diagnostic messages regarding changes in contact status selectively only when a detailed contact printout is requested.

### 1.1.36 UMASFL: User subroutine to specify prescribed mass flow rate conditions for a convection/diffusion heat transfer analysis.

**Product:** Abaqus/Standard

#### References

---

- “Uncoupled heat transfer analysis,” Section 6.5.2 of the Abaqus Analysis User’s Manual
- \*MASS FLOW RATE
- “UTEMP, UFIELD, UMASFL, and UPRESS,” Section 4.1.25 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UMASFL**:

- can be used to prescribe the mass flow rate vector at the nodes of a model as a function of position and time;
- will be called whenever a current value of mass flow rate (per unit area) is needed for a node listed in a user-subroutine-defined mass flow rate definition (the node should belong to one or more convection/diffusion elements); and
- will overwrite any flow rate data specified for the associated mass flow rate definition outside the user subroutine.

#### User subroutine interface

---

```

SUBROUTINE UMASFL (FLOW, KFLOW, KSTEP, KINC, TIME, NODE, COORDS)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION FLOW(KFLOW), TIME(2), COORDS(3)
C

```

*user coding to define FLOW*

```

RETURN
END

```

---

**Variable to be defined****FLOW**

Total value of the mass flow rate vector at this point. The number of components in this vector is **KFLOW**. If **KFLOW**=1, give the total mass flow rate through the cross-section (for one-dimensional elements). If **KFLOW**=2, give the *x*-component and *y*-component of the flow rate vector as **FLOW(1)** and **FLOW(2)**. If **KFLOW**=3, give the *x*-component, *y*-component, and *z*-component as **FLOW(1)**, **FLOW(2)**, and **FLOW(3)**.

---

**Variables passed in for information****KFLOW**

The number of components in the mass flow rate vector. **UMASFL** will be called only once per node.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME(1)**

Current value of step time.

**TIME(2)**

Current value of total time.

**NODE**

Node number.

**COORDS**

An array containing the coordinates of this node.

### 1.1.37 UMAT: User subroutine to define a material's mechanical behavior.

**Product:** Abaqus/Standard

*WARNING: The use of this subroutine generally requires considerable expertise. You are cautioned that the implementation of any realistic constitutive model requires extensive development and testing. Initial testing on a single-element model with prescribed traction loading is strongly recommended.*

#### References

---

- “User-defined mechanical material behavior,” Section 23.8.1 of the Abaqus Analysis User’s Manual
- “User-defined thermal material behavior,” Section 23.8.2 of the Abaqus Analysis User’s Manual
- \*USER MATERIAL
- “**SDVINI**,” Section 4.1.11 of the Abaqus Verification Manual
- “**UMAT** and **UHYPER**,” Section 4.1.21 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UMAT**:

- can be used to define the mechanical constitutive behavior of a material;
- will be called at all material calculation points of elements for which the material definition includes a user-defined material behavior;
- can be used with any procedure that includes mechanical behavior;
- can use solution-dependent state variables;
- must update the stresses and solution-dependent state variables to their values at the end of the increment for which it is called;
- must provide the material Jacobian matrix,  $\partial\Delta\sigma/\partial\Delta\varepsilon$ , for the mechanical constitutive model;
- can be used in conjunction with user subroutine **USDFLD** to redefine any field variables before they are passed in; and
- is described further in “User-defined mechanical material behavior,” Section 23.8.1 of the Abaqus Analysis User’s Manual.

#### Storage of stress and strain components

---

In the stress and strain arrays and in the matrices **DDSDDE**, **DDSDDT**, and **DRPLDE**, direct components are stored first, followed by shear components. There are **NDI** direct and **NSHR** engineering shear components. The order of the components is defined in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual. Since the number of active stress and strain components varies between element types, the routine must be coded to provide for all element types with which it will be used.

## Defining local orientations

---

If a local orientation (“Orientations,” Section 2.2.5 of the Abaqus Analysis User’s Manual) is used at the same point as user subroutine **UMAT**, the stress and strain components will be in the local orientation; and, in the case of finite-strain analysis, the basis system in which stress and strain components are stored rotates with the material.

## Stability

---

You should ensure that the integration scheme coded in this routine is stable—no direct provision is made to include a stability limit in the time stepping scheme based on the calculations in **UMAT**.

## Convergence rate

---

**DDSDDE** and, for coupled temperature-displacement analyses, **DDSDDT**, **DRPLDE**, and **DRPLDT**, must be defined accurately if rapid convergence of the overall Newton scheme is to be achieved. In most cases the accuracy of this definition is the most important factor governing the convergence rate. Since nonsymmetric equation solution is as much as four times as expensive as the corresponding symmetric system, if the constitutive Jacobian (**DDSDDE**) is only slightly nonsymmetric (for example, a frictional material with a small friction angle), it may be less expensive computationally to use a symmetric approximation and accept a slower convergence rate.

An incorrect definition of the material Jacobian affects only the convergence rate; the results (if obtained) are unaffected.

## Special considerations for various element types

---

There are several special considerations that need to be noted.

### Availability of deformation gradient

The deformation gradient is available for solid (continuum) elements, membranes, and finite-strain shells (S3/S3R, S4, S4R, SAXs, and SAXAs). It is not available for beams or small-strain shells. It is stored as a  $3 \times 3$  matrix with component equivalence  $\text{DFGRD0}(\mathbf{I}, \mathbf{J}) \Leftrightarrow F_{I,J}$ . For fully integrated first-order isoparametric elements (4-node quadrilaterals in two dimensions and 8-node hexahedra in three dimensions) the selectively reduced integration technique is used (also known as the  $\bar{B}$  technique). Thus, a modified deformation gradient

$$\bar{\mathbf{F}} = \mathbf{F} \left( \frac{\bar{J}}{J} \right)^{\frac{1}{n}}$$

is passed into user subroutine **UMAT**. For more details, see “Solid isoparametric quadrilaterals and hexahedra,” Section 3.2.4 of the Abaqus Theory Manual.

## Beams and shells that calculate transverse shear energy

If user subroutine **UMAT** is used to describe the material of beams or shells that calculate transverse shear energy, you must specify the transverse shear stiffness as part of the beam or shell section definition to define the transverse shear behavior. See “Shell section behavior,” Section 26.6.4 of the Abaqus Analysis User’s Manual, and “Choosing a beam element,” Section 26.3.3 of the Abaqus Analysis User’s Manual, for information on specifying this stiffness.

## Open-section beam elements

When user subroutine **UMAT** is used to describe the material response of beams with open sections (for example, an I-section), the torsional stiffness is obtained as

$$GJ = \frac{(K_{13} + K_{23})J}{2kA},$$

where  $J$  is the torsional constant,  $A$  is the section area,  $k$  is a shear factor, and  $K_{\alpha 3}$  is the user-specified transverse shear stiffness (see “Transverse shear stiffness definition” in “Choosing a beam element,” Section 26.3.3 of the Abaqus Analysis User’s Manual).

## Elements with hourglassing modes

If this capability is used to describe the material of elements with hourglassing modes, you must define the hourglass stiffness factor for hourglass control based on the total stiffness approach as part of the element section definition. The hourglass stiffness factor is not required for enhanced hourglass control, but you can define a scaling factor for the stiffness associated with the drill degree of freedom (rotation about the surface normal). See “Section controls,” Section 24.1.4 of the Abaqus Analysis User’s Manual, for information on specifying the stiffness factor.

## Pipe-soil interaction elements

The constitutive behavior of the pipe-soil interaction elements (see “Pipe-soil interaction elements,” Section 29.13.1 of the Abaqus Analysis User’s Manual) is defined by the force per unit length caused by relative displacement between two edges of the element. The relative-displacements are available as “strains” (**STRAN** and **DSTRAN**). The corresponding forces per unit length must be defined in the **STRESS** array. The Jacobian matrix defines the variation of force per unit length with respect to relative displacement.

For two-dimensional elements two in-plane components of “stress” and “strain” exist (**NTENS=NDI=2**, and **NSHR=0**). For three-dimensional elements three components of “stress” and “strain” exist (**NTENS=NDI=3**, and **NSHR=0**).

## Large volume changes with geometric nonlinearity

If the material model allows large volume changes and geometric nonlinearity is considered, the exact definition of the consistent Jacobian should be used to ensure rapid convergence. These conditions are most commonly encountered when considering either large elastic strains or pressure-dependent

plasticity. In the former case, total-form constitutive equations relating the Cauchy stress to the deformation gradient are commonly used; in the latter case, rate-form constitutive laws are generally used.

For total-form constitutive laws, the exact consistent Jacobian  $\mathbf{C}$  is defined through the variation in Kirchhoff stress:

$$\delta(J\sigma) = J(\mathbf{C} : \delta\mathbf{D} + \delta\mathbf{W} \cdot \boldsymbol{\sigma} - \boldsymbol{\sigma} \cdot \delta\mathbf{W})$$

Here,  $J$  is the determinant of the deformation gradient,  $\boldsymbol{\sigma}$  is the Cauchy stress,  $\delta\mathbf{D}$  is the virtual rate of deformation, and  $\delta\mathbf{W}$  is the virtual spin tensor, defined as

$$\delta\mathbf{D} \stackrel{\text{def}}{=} \text{sym}(\delta\mathbf{F} \cdot \mathbf{F}^{-1})$$

and

$$\delta\mathbf{W} \stackrel{\text{def}}{=} \text{asym}(\delta\mathbf{F} \cdot \mathbf{F}^{-1}).$$

For rate-form constitutive laws, the exact consistent Jacobian is given by

$$\mathbf{C} = \frac{1}{J} \frac{\partial \Delta(J\sigma)}{\partial \Delta\epsilon}.$$

## **Use with incompressible elastic materials**

---

For user-defined incompressible elastic materials, user subroutine **UHYPER** should be used rather than user subroutine **UMAT**. In **UMAT** incompressible materials must be modeled via a penalty method; that is, you must ensure that a finite bulk modulus is used. The bulk modulus should be large enough to model incompressibility sufficiently but small enough to avoid loss of precision. As a general guideline, the bulk modulus should be about  $10^4$ – $10^6$  times the shear modulus. The tangent bulk modulus  $K^t$  can be calculated from

$$K^t = \frac{1}{9} \sum_{I=1}^3 \sum_{J=1}^3 \text{DDSDDE}(I, J).$$

If a hybrid element is used with user subroutine **UMAT**, Abaqus/Standard will replace the pressure stress calculated from your definition of **STRESS** with that derived from the Lagrange multiplier and will modify the Jacobian appropriately.

For incompressible pressure-sensitive materials the element choice is particularly important when using user subroutine **UMAT**. In particular, first-order wedge elements should be avoided. For these elements the  $\bar{B}$  technique is not used to alter the deformation gradient that is passed into user subroutine **UMAT**, which increases the risk of volumetric locking.

---

## Increments for which only the Jacobian can be defined

---

Abaqus/Standard passes zero strain increments into user subroutine **UMAT** to start the first increment of all the steps and all increments of steps for which you have suppressed extrapolation (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual). In this case you can define only the Jacobian (**DDSDDE**).

---

## Utility routines

---

Several utility routines may help in coding user subroutine **UMAT**. Their functions include determining stress invariants for a stress tensor and calculating principal values and directions for stress or strain tensors. These utility routines are discussed in detail in “Obtaining stress invariants, principal stress/strain values and directions, and rotating tensors in an Abaqus/Standard analysis,” Section 2.1.11.

---

## User subroutine interface

---

```
SUBROUTINE UMAT(STRESS,STATEV,DDSDDE,SSE,SPD,SCD,
  1 RPL,DDSDDT,DRPLDE,DRPLDT,
  2 STRAN,DSTRAN,TIME,DTIME,TEMP,DTEMP,PREDEF,DPRED,CMNAME,
  3 NDI,NSHR,NTENS,NSTATV,PROPS,NPROPS,COORDS,DROT,PNEWDT,
  4 CELENT,DFGRD0,DFGRD1,NOEL,NPT,LAYER,KSPT,KSTEP,KINC)
C
  INCLUDE 'ABA_PARAM.INC'
C
  CHARACTER*80 CMNAME
  DIMENSION STRESS(NTENS),STATEV(NSTATV),
  1 DDSDDE(NTENS,NTENS),DDSDDT(NTENS),DRPLDE(NTENS),
  2 STRAN(NTENS),DSTRAN(NTENS),TIME(2),PREDEF(1),DPRED(1),
  3 PROPS(NPROPS),COORDS(3),DROT(3,3),DFGRD0(3,3),DFGRD1(3,3)
```

*user coding to define DDSDDE, STRESS, STATEV, SSE, SPD, SCD  
and, if necessary, RPL, DDSDDT, DRPLDE, DRPLDT, PNEWDT*

```
RETURN
END
```

---

## Variables to be defined

---

### In all situations

#### **DDSDDE (NTENS, NTENS)**

Jacobian matrix of the constitutive model,  $\partial\Delta\sigma/\partial\Delta\varepsilon$ , where  $\Delta\sigma$  are the stress increments and  $\Delta\varepsilon$  are the strain increments. **DDSDDE (I, J)** defines the change in the Ith stress component at the end of the

time increment caused by an infinitesimal perturbation of the **J**th component of the strain increment array. Unless you invoke the unsymmetric equation solution capability for the user-defined material, Abaqus/Standard will use only the symmetric part of **DDSDDE**. The symmetric part of the matrix is calculated by taking one half the sum of the matrix and its transpose.

#### **STRESS (NTENS)**

This array is passed in as the stress tensor at the beginning of the increment and must be updated in this routine to be the stress tensor at the end of the increment. If you specified initial stresses (“Initial conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.2.1 of the Abaqus Analysis User’s Manual), this array will contain the initial stresses at the start of the analysis. The size of this array depends on the value of **NTENS** as defined below. In finite-strain problems the stress tensor has already been rotated to account for rigid body motion in the increment before **UMAT** is called, so that only the corotational part of the stress integration should be done in **UMAT**. The measure of stress used is “true” (Cauchy) stress.

#### **STATEV (NSTATV)**

An array containing the solution-dependent state variables. These are passed in as the values at the beginning of the increment unless they are updated in user subroutines **USDFLD** or **UEXPAN**, in which case the updated values are passed in. In all cases **STATEV** must be returned as the values at the end of the increment. The size of the array is defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

In finite-strain problems any vector-valued or tensor-valued state variables must be rotated to account for rigid body motion of the material, in addition to any update in the values associated with constitutive behavior. The rotation increment matrix, **DROT**, is provided for this purpose.

#### **SSE, SPD, SCD**

Specific elastic strain energy, plastic dissipation, and “creep” dissipation, respectively. These are passed in as the values at the start of the increment and should be updated to the corresponding specific energy values at the end of the increment. They have no effect on the solution, except that they are used for energy output.

#### **Only in a fully coupled thermal-stress analysis**

##### **RPL**

Volumetric heat generation per unit time at the end of the increment caused by mechanical working of the material.

##### **DDSDDT (NTENS)**

Variation of the stress increments with respect to the temperature.

##### **DRPLDE (NTENS)**

Variation of **RPL** with respect to the strain increments.

##### **DRPLDT**

Variation of **RPL** with respect to the temperature.

## Variable that can be updated

---

### **PNEWDT**

Ratio of suggested new time increment to the time increment being used (**DTIME**, see discussion later in this section). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen). For a quasi-static procedure the automatic time stepping that Abaqus/Standard uses, which is based on techniques for integrating standard creep laws (see “Quasi-static analysis,” Section 6.2.5 of the Abaqus Analysis User’s Manual), cannot be controlled from within the **UMAT** subroutine.

**PNEWDT** is set to a large value before each call to **UMAT**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is  $\text{PNEWDT} \times \text{DTIME}$ , where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is  $\text{PNEWDT} \times \text{DTIME}$ , where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

## Variables passed in for information

---

### **STRAN (NTENS)**

An array containing the total strains at the beginning of the increment. If thermal expansion is included in the same material definition, the strains passed into **UMAT** are the mechanical strains only (that is, the thermal strains computed based upon the thermal expansion coefficient have been subtracted from the total strains). These strains are available for output as the “elastic” strains.

In finite-strain problems the strain components have been rotated to account for rigid body motion in the increment before **UMAT** is called and are approximations to logarithmic strain.

### **DSTRAN (NTENS)**

Array of strain increments. If thermal expansion is included in the same material definition, these are the mechanical strain increments (the total strain increments minus the thermal strain increments).

### **TIME (1)**

Value of step time at the beginning of the current increment.

### **TIME (2)**

Value of total time at the beginning of the current increment.

### **DTIME**

Time increment.

**TEMP**

Temperature at the start of the increment.

**DTEMP**

Increment of temperature.

**PREDEF**

Array of interpolated values of predefined field variables at this point at the start of the increment, based on the values read in at the nodes.

**DPRED**

Array of increments of predefined field variables.

**CMNAME**

User-defined material name, left justified. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, you should not use “ABQ\_” as the leading string for **CMNAME**.

**NDI**

Number of direct stress components at this point.

**NSHR**

Number of engineering shear stress components at this point.

**NTENS**

Size of the stress or strain component array (**NDI** + **NSHR**).

**NSTATV**

Number of solution-dependent state variables that are associated with this material type (defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**PROPS (NPROPS)**

User-specified array of material constants associated with this user material.

**NPROPS**

User-defined number of material constants associated with this user material.

**COORDS**

An array containing the coordinates of this point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**DRROT (3 , 3)**

Rotation increment matrix. This matrix represents the increment of rigid body rotation of the basis system in which the components of stress (**STRESS**) and strain (**STRAN**) are stored. It is provided so

that vector- or tensor-valued state variables can be rotated appropriately in this subroutine: stress and strain components are already rotated by this amount before **UMAT** is called. This matrix is passed in as a unit matrix for small-displacement analysis and for large-displacement analysis if the basis system for the material point rotates with the material (as in a shell element or when a local orientation is used).

#### **CELENT**

Characteristic element length, which is a typical length of a line across an element for a first-order element; it is half of the same typical length for a second-order element. For beams and trusses it is a characteristic length along the element axis. For membranes and shells it is a characteristic length in the reference surface. For axisymmetric elements it is a characteristic length in the  $(r, z)$  plane only. For cohesive elements it is equal to the constitutive thickness.

#### **DFGRD0 (3, 3)**

Array containing the deformation gradient at the beginning of the increment. For a discussion regarding the availability of the deformation gradient for various element types, see “Availability of deformation gradient.”

#### **DFGRD1 (3, 3)**

Array containing the deformation gradient at the end of the increment. This array is set to the identity matrix if nonlinear geometric effects are not included in the step definition associated with this increment. For a discussion regarding the availability of the deformation gradient for various element types, see “Availability of deformation gradient.”

#### **NOEL**

Element number.

#### **NPT**

Integration point number.

#### **LAYER**

Layer number (for composite shells and layered solids).

#### **KSPT**

Section point number within the current layer.

#### **KSTEP**

Step number.

#### **KINC**

Increment number.

---

#### **Example: Using more than one user-defined mechanical material model**

To use more than one user-defined mechanical material model, the variable **CMMNAME** can be tested for different material names inside user subroutine **UMAT** as illustrated below:

```

IF (CMNAME(1:4) .EQ. 'MAT1') THEN
  CALL UMAT_MAT1(argument_list)
ELSE IF(CMNAME(1:4) .EQ. 'MAT2') THEN
  CALL UMAT_MAT2(argument_list)
END IF

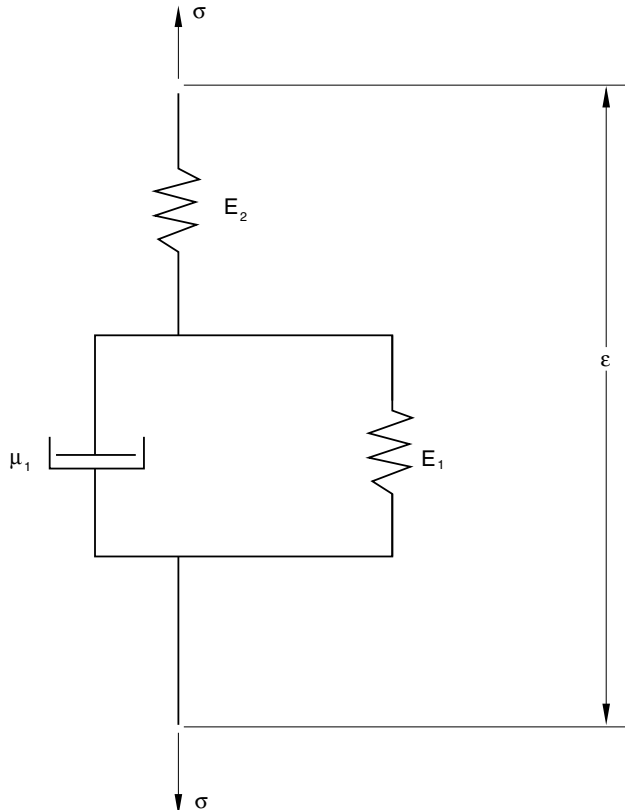
```

**UMAT\_MAT1** and **UMAT\_MAT2** are the actual user material subroutines containing the constitutive material models for each material **MAT1** and **MAT2**, respectively. Subroutine **UMAT** merely acts as a directory here. The argument list may be the same as that used in subroutine **UMAT**.

#### Example: Simple linear viscoelastic material

---

As a simple example of the coding of user subroutine **UMAT**, consider the linear, viscoelastic model shown in Figure 1.1.37–1. Although this is not a very useful model for real materials, it serves to illustrate how to code the routine.



**Figure 1.1.37–1** Simple linear viscoelastic model.

The behavior of the one-dimensional model shown in the figure is

$$\sigma + \frac{\mu_1}{(E_1 + E_2)}\dot{\sigma} = \frac{\mu_1}{(1 + E_1/E_2)}\dot{\varepsilon} + \frac{1}{(1/E_1 + 1/E_2)}\varepsilon,$$

where  $\dot{\sigma}$  and  $\dot{\varepsilon}$  are the time rates of change of stress and strain. This can be generalized for small straining of an isotropic solid as

$$\sigma_{xx} + \tilde{\nu}\dot{\sigma}_{xx} = \lambda\varepsilon_V + 2\mu\varepsilon_{xx} + \tilde{\lambda}\dot{\varepsilon}_V + 2\tilde{\mu}\dot{\varepsilon}_{xx}, \quad \text{etc.,}$$

and

$$\sigma_{xy} + \tilde{\nu}\dot{\sigma}_{xy} = \mu\gamma_{xy} + \tilde{\mu}\dot{\gamma}_{xy}, \quad \text{etc.,}$$

where

$$\varepsilon_V = \varepsilon_{xx} + \varepsilon_{yy} + \varepsilon_{zz},$$

and  $\tilde{\nu}$ ,  $\lambda$ ,  $\mu$ ,  $\tilde{\lambda}$ , and  $\tilde{\mu}$  are material constants ( $\lambda$  and  $\mu$  are the Lamé constants).

A simple, stable integration operator for this equation is the central difference operator:

$$\begin{aligned}\dot{f}_{t+\frac{1}{2}\Delta t} &= \frac{\Delta f}{\Delta t}, \\ f_{t+\frac{1}{2}\Delta t} &= f_t + \frac{\Delta f}{2},\end{aligned}$$

where  $f$  is some function,  $f_t$  is its value at the beginning of the increment,  $\Delta f$  is the change in the function over the increment, and  $\Delta t$  is the time increment.

Applying this to the rate constitutive equations above gives

$$(\frac{\Delta t}{2} + \tilde{\nu})\Delta\sigma_{xx} = (\Delta t\frac{\lambda}{2} + \tilde{\lambda})\Delta\varepsilon_V + (\Delta t\mu + 2\tilde{\mu})\Delta\varepsilon_{xx} + \Delta t(\lambda\varepsilon_V + 2\mu\varepsilon_{xx} - \sigma_{xx})_t, \quad \text{etc.,}$$

and

$$(\frac{\Delta t}{2} + \tilde{\nu})\Delta\sigma_{xy} = (\Delta t\frac{\mu}{2} + \tilde{\mu})\Delta\gamma_{xy} + \Delta t(\mu\gamma_{xy} - \sigma_{xy})_t, \quad \text{etc.,}$$

so that the Jacobian matrix has the terms

$$\frac{\partial\Delta\sigma_{xx}}{\partial\Delta\varepsilon_{xx}} = \frac{1}{(\frac{\Delta t}{2} + \tilde{\nu})}[\Delta t(\frac{\lambda}{2} + \mu) + \tilde{\lambda} + 2\tilde{\mu}],$$

$$\frac{\partial\Delta\sigma_{xx}}{\partial\Delta\varepsilon_{yy}} = \frac{1}{(\frac{\Delta t}{2} + \tilde{\nu})}[\Delta t\frac{\lambda}{2} + \tilde{\lambda}],$$

and

$$\frac{\partial \Delta\sigma_{xy}}{\partial \Delta\gamma_{xy}} = \frac{1}{(\frac{\Delta t}{2} + \tilde{\nu})} [\Delta t \frac{\mu}{2} + \tilde{\mu}].$$

The total change in specific energy in an increment for this material is

$$(\sigma_{ij} + \frac{1}{2}\Delta\sigma_{ij})\Delta\varepsilon_{ij},$$

while the change in specific elastic strain energy is

$$(\varepsilon_{ij} + \frac{1}{2}\Delta\varepsilon_{ij})D_{ijkl}\Delta\varepsilon_{kl},$$

where  $D$  is the elasticity matrix:

$$\begin{bmatrix} \lambda + 2\mu & \lambda & \lambda & 0 & 0 & 0 \\ \lambda & \lambda + 2\mu & \lambda & 0 & 0 & 0 \\ \lambda & \lambda & \lambda + 2\mu & 0 & 0 & 0 \\ 0 & 0 & 0 & \mu & 0 & 0 \\ 0 & 0 & 0 & 0 & \mu & 0 \\ 0 & 0 & 0 & 0 & 0 & \mu \end{bmatrix}.$$

No state variables are needed for this material, so the allocation of space for them is not necessary. In a more realistic case a set of parallel models of this type might be used, and the stress components in each model might be stored as state variables.

For our simple case a user material definition can be used to read in the five constants in the order  $\lambda, \mu, \tilde{\lambda}, \tilde{\mu}$ , and  $\tilde{\nu}$  so that

$$\text{PROPS}(1) = \lambda,$$

$$\text{PROPS}(2) = \mu,$$

$$\text{PROPS}(3) = \tilde{\lambda},$$

$$\text{PROPS}(4) = \tilde{\mu},$$

$$\text{PROPS}(5) = \tilde{\nu}.$$

The routine can then be coded as follows:

```
SUBROUTINE UMAT(STRESS,STATEV,DDSDDE,SSE,SPD,SCD,
1 RPL,DDSDDT,DRPLDE,DRPLDT,
```

```

2 STRAN,DSTRAN,TIME,DTIME,TEMP,DTEMP,PREDEF,DPRED,CMNAME,
3 NDI,NSHR,NTENS,NSTATV,PROPS,NPROPS,COORDS,DROT,PNEWDT,
4 CELENT,DFGRD0,DFGRD1,NOEL,NPT,LAYER,KSPT,KSTEP,KINC)
C
C INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
DIMENSION STRESS(NTENS),STATEV(NSTATV),
1 DDSdde(NTENS,NTENS),
2 DDSddt(NTENS),DRPLDE(NTENS),
3 STRAN(NTENS),DSTRAN(NTENS),TIME(2),PREDEF(1),DPRED(1),
4 PROPS(NPROPS),COORDS(3),DROT(3,3),DFGRD0(3,3),DFGRD1(3,3)
DIMENSION DSTRES(6),D(3,3)

C
C EVALUATE NEW STRESS TENSOR
C
EV = 0.
DEV = 0.
DO K1=1,NDI
    EV = EV + STRAN(K1)
    DEV = DEV + DSTRAN(K1)
END DO
C
TERM1 = .5*DTIME + PROPS(5)
TERM1I = 1./TERM1
TERM2 = (.5*DTIME*PROPS(1)+PROPS(3))*TERM1I*DEV
TERM3 = (DTIME*PROPS(2)+2.*PROPS(4))*TERM1I
C
DO K1=1,NDI
    DSTRES(K1) = TERM2+TERM3*DSTRAN(K1)
1     +DTIME*TERM1I*(PROPS(1)*EV
2     +2.*PROPS(2)*STRAN(K1)-STRESS(K1))
    STRESS(K1) = STRESS(K1) + DSTRES(K1)
END DO
C
TERM2 = (.5*DTIME*PROPS(2) + PROPS(4))*TERM1I
I1 = NDI
DO K1=1,NSHR
    I1 = I1+1
    DSTRES(I1) = TERM2*DSTRAN(I1) +
1     DTIME*TERM1I*(PROPS(2)*STRAN(I1)-STRESS(I1))
    STRESS(I1) = STRESS(I1)+DSTRES(I1)

```

```

        END DO
C
C  CREATE NEW JACOBIAN
C
    TERM2 = (DTIME*(.5*PROPS(1)+PROPS(2))+PROPS(3) +
1  2.*PROPS(4))*TERM1I
    TERM3 = (.5*DTIME*PROPS(1)+PROPS(3))*TERM1I
    DO K1=1,NTENS
        DO K2=1,NTENS
            DDSDDE(K2,K1) = 0.
        END DO
    END DO
C
    DO K1=1,NDI
        DDSDDE(K1,K1) = TERM2
    END DO
C
    DO K1=2,NDI
        N2 = K1-1
        DO K2=1,N2
            DDSDDE(K2,K1) = TERM3
            DDSDDE(K1,K2) = TERM3
        END DO
    END DO
    TERM2 = (.5*DTIME*PROPS(2)+PROPS(4))*TERM1I
    I1 = NDI
    DO K1=1,NSHR
        I1 = I1+1
        DDSDDE(I1,I1) = TERM2
    END DO
C
C  TOTAL CHANGE IN SPECIFIC ENERGY
C
    TDE = 0.
    DO K1=1,NTENS
        TDE = TDE + (STRESS(K1)-.5*DSTRES(K1))*DSTRAN(K1)
    END DO
C
C  CHANGE IN SPECIFIC ELASTIC STRAIN ENERGY
C
    TERM1 = PROPS(1) + 2.*PROPS(2)
    DO K1=1,NDI

```

```
D (K1 ,K1) = TERM1
END DO
DO K1=2,NDI
    N2 = K1-1
    DO K2=1,N2
        D (K1 ,K2) = PROPS (1)
        D (K2 ,K1) = PROPS (1)
    END DO
END DO
DEE = 0 .
DO K1=1,NDI
    TERM1 = 0 .
    TERM2 = 0 .
    DO K2=1,NDI
        TERM1 = TERM1 + D (K1 ,K2) *STRAN (K2)
        TERM2 = TERM2 + D (K1 ,K2) *DSTRAN (K2)
    END DO
    DEE = DEE + (TERM1+.5*TERM2) *DSTRAN (K1)
END DO
I1 = NDI
DO K1=1,NSHR
    I1 = I1+1
    DEE = DEE + PROPS (2) *(STRAN (I1)+.5*DSTRAN (I1)) *DSTRAN (I1)
END DO
SSE = SSE + DEE
SCD = SCD + TDE - DEE
RETURN
END
```



### 1.1.38 UMATHT: User subroutine to define a material's thermal behavior.

**Product:** Abaqus/Standard

*WARNING: The use of this subroutine generally requires considerable expertise. You are cautioned that the implementation of any realistic thermal model requires significant development and testing. Initial testing on models with few elements under a variety of boundary conditions is strongly recommended.*

#### References

---

- “User-defined thermal material behavior,” Section 23.8.2 of the Abaqus Analysis User’s Manual
- \*USER MATERIAL
- “Freezing of a square solid: the two-dimensional Stefan problem,” Section 1.6.2 of the Abaqus Benchmarks Manual
- “UMATHT,” Section 4.1.22 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UMATHT**:

- can be used to define the thermal constitutive behavior of the material as well as internal heat generation during heat transfer processes;
- will be called at all material calculation points of elements for which the material definition includes a user-defined thermal material behavior;
- can be used with the procedures discussed in “Heat transfer analysis procedures: overview,” Section 6.5.1 of the Abaqus Analysis User’s Manual;
- can use solution-dependent state variables;
- must define the internal energy per unit mass and its variation with respect to temperature and to spatial gradients of temperature;
- must define the heat flux vector and its variation with respect to temperature and to gradients of temperature;
- must update the solution-dependent state variables to their values at the end of the increment;
- can be used in conjunction with user subroutine **USDFLD** to redefine any field variables before they are passed in; and
- is described further in “User-defined thermal material behavior,” Section 23.8.2 of the Abaqus Analysis User’s Manual.

## Use of subroutine UMATHT with coupled temperature-displacement elements

---

User subroutine **UMATHT** should be used only with reduced-integration or modified coupled temperature-displacement elements if the mechanical and thermal fields are not coupled through plastic dissipation. No such restriction exists with fully integrated coupled temperature-displacement elements.

## User subroutine interface

---

```

SUBROUTINE UMATHT (U,DUDT,DUDG,FLUX,DFDT,DFDG,
 1 STATEV,TEMP,DTEMP,DTEMDX,TIME,DTIME,PREDEF,DPRED,
 2 CMNAME,NTGRD,NSTATV,PROPS,NPROPS,COORDS,PNEWDT,
 3 NOEL,NPT,LAYER,KSPT,KSTEP,KINC)
C
  INCLUDE 'ABA_PARAM.INC'
C
  CHARACTER*80 CMNAME
  DIMENSION DUDG(NTGRD),FLUX(NTGRD),DFDT(NTGRD),
 1 DFDG(NTGRD,NTGRD),STATEV(NSTATV),DTEMDX(NTGRD),
 2 TIME(2),PREDEF(1),DPRED(1),PROPS(NPROPS),COORDS(3)

```

*user coding to define **U**, **DUDT**, **DUDG**, **FLUX**, **DFDT**, **DFDG**,  
and possibly update **STATEV**, **PNEWDT***

```

RETURN
END

```

## Variables to be defined

---

**U**

Internal thermal energy per unit mass,  $U$ , at the end of increment. This variable is passed in as the value at the start of the increment and must be updated to its value at the end of the increment.

**DUDT**

Variation of internal thermal energy per unit mass with respect to temperature,  $\partial U / \partial \theta$ , evaluated at the end of the increment.

**DUDG (NTGRD)**

Variation of internal thermal energy per unit mass with respect to the spatial gradients of temperature,  $\partial U / \partial (\partial \theta / \partial x)$ , at the end of the increment. The size of this array depends on the value of **NTGRD** as defined below. This term is typically zero in classical heat transfer analysis.

**FLUX (NTGRD)**

Heat flux vector,  $\mathbf{f}$ , at the end of the increment. This variable is passed in with the values at the beginning of the increment and must be updated to the values at the end of the increment.

**DFDT (NTGRD)**

Variation of the heat flux vector with respect to temperature,  $\partial\mathbf{f}/\partial\theta$ , evaluated at the end of the increment.

**DFDG (NTGRD , NTGRD)**

Variation of the heat flux vector with respect to the spatial gradients of temperature,  $\partial\mathbf{f}/\partial(\partial\theta/\partial\mathbf{x})$ , at the end of the increment. The size of this array depends on the value of **NTGRD** as defined below.

**Variables that can be updated**

---

**STATEV (NSTATV)**

An array containing the solution-dependent state variables.

In an uncoupled heat transfer analysis **STATEV** is passed into **UMATHT** with the values of these variables at the beginning of the increment. However, any changes in **STATEV** made in user subroutine **USDFLD** will be included in the values passed into **UMATHT**, since **USDFLD** is called before **UMATHT**. In addition, if **UMATHT** is being used in a fully coupled temperature-displacement analysis and user subroutine **CREEP**, user subroutine **UEXPAN**, user subroutine **UMAT**, or user subroutine **UTRS** is used to define the mechanical behavior of the material, those routines are called before this routine; therefore, any updating of **STATEV** done in **CREEP**, **UEXPAN**, **UMAT**, or **UTRS** will be included in the values passed into **UMATHT**.

In all cases **STATEV** should be passed back from **UMATHT** as the values of the state variables at the end of the current increment.

**PNEWDT**

Ratio of suggested new time increment to the time increment being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **UMATHT**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT**  $\times$  **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT**  $\times$  **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

---

**Variables passed in for information****TEMP**

Temperature at the start of the increment.

**DTEMP**

Increment of temperature.

**DTEMDX (NTGRD)**

Current values of the spatial gradients of temperature,  $\partial\theta/\partial\mathbf{x}$ .

**TIME (1)**

Value of step time at the beginning of the current increment.

**TIME (2)**

Value of total time at the beginning of the current increment.

**DTIME**

Time increment.

**PREFDEF**

Array of interpolated values of predefined field variables at this point at the start of the increment, based on the values read in at the nodes.

**DPRED**

Array of increments of predefined field variables.

**CMNAME**

User-defined material name, left justified.

**NTGRD**

Number of spatial gradients of temperature.

**NSTATTV**

Number of solution-dependent state variables associated with this material type (defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**PROPS (NPROPS)**

User-specified array of material constants associated with this user material.

**NPROPS**

User-defined number of material constants associated with this user material.

**COORDS**

An array containing the coordinates of this point. These are the current coordinates in a fully coupled temperature-displacement analysis if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**KSTEP**

Step number.

**KINC**

Increment number.

**Example: Using more than one user-defined thermal material model**

---

To use more than one user-defined thermal material model, the variable **CMNAME** can be tested for different material names inside user subroutine **UMATHT**, as illustrated below:

```
IF (CMNAME(1:4) .EQ. 'MAT1') THEN
    CALL UMATHT_MAT1(argument_list)
ELSE IF(CMNAME(1:4) .EQ. 'MAT2') THEN
    CALL UMATHT_MAT2(argument_list)
END IF
```

**UMATHT\_MAT1** and **UMATHT\_MAT2** are the actual user material subroutines containing the constitutive material models for each material **MAT1** and **MAT2**, respectively. Subroutine **UMATHT** merely acts as a directory here. The argument list can be the same as that used in subroutine **UMATHT**.

**Example: Uncoupled heat transfer**

---

As a simple example of the coding of user subroutine **UMATHT**, consider uncoupled heat transfer analysis in a material. The equations for this case are developed here, and the corresponding **UMATHT** is given. This problem can also be solved by specifying thermal conductivity, specific heat, density, and internal heat generation directly.

First, the equations for an uncoupled heat transfer analysis are outlined.  
The basic energy balance is

$$\int_V \rho \dot{U} dV = \int_S q dS + \int_V r dV,$$

where  $V$  is the volume of solid material with surface area  $S$ ,  $\rho$  is the density of the material,  $\dot{U}$  is the material time rate of the internal thermal energy,  $q$  is the heat flux per unit area of the body flowing into the body, and  $r$  is the heat supplied externally into the body per unit volume.

A heat flux vector  $\mathbf{f}$  is defined such that

$$q = -\mathbf{f} \cdot \mathbf{n},$$

where  $\mathbf{n}$  is the unit outward normal to the surface  $S$ . Introducing the above relation into the energy balance equation and using the divergence theorem, the following relation is obtained:

$$\int_V \rho \dot{U} dV = - \int_V \frac{\partial}{\partial \mathbf{x}} \cdot \mathbf{f} dV + \int_V r dV.$$

The corresponding weak form is given by

$$\int_V \delta \theta \rho \dot{U} dV - \int_V \delta \mathbf{g} \cdot \mathbf{f} dV = \int_V \delta \theta r dV + \int_S \delta \theta q dS,$$

where

$$\mathbf{g} = \frac{\partial \theta}{\partial \mathbf{x}}$$

is the temperature gradient and  $\delta \theta$  is an arbitrary variational field satisfying the essential boundary conditions.

Introducing the backward difference integration algorithm:

$$\dot{U}_{t+\Delta t} = (U_{t+\Delta t} - U_t)(1/\Delta t),$$

the weak form of the energy balance equation becomes

$$\frac{1}{\Delta t} \int_V \delta \theta \rho (U_{t+\Delta t} - U_t) dV = \int_V \delta \mathbf{g} \cdot \mathbf{f} dV + \int_V \delta \theta r dV + \int_S \delta \theta q dS.$$

This nonlinear system is solved using Newton's method.

In the above equations the thermal constitutive behavior of the material is given by

$$U = U(\theta, t, \partial \theta / \partial \mathbf{x}, s^i, \dots) U = U(\theta, \mathbf{g}, t, s^i, \dots)$$

and

$$\mathbf{f} = \mathbf{f}(\theta, t, \partial\theta/\partial\mathbf{x}, s^i, \dots), \mathbf{f} = \mathbf{f}(\theta, \mathbf{g}, t, s^i, \dots),$$

where  $s^i$  are state variables.

The Jacobian for Newton's method is given by (after dropping the subscripts  $t + \Delta t$  on  $U$ )

$$\begin{aligned} \frac{1}{\Delta t} \int_V \delta\theta \rho \frac{\partial U}{\partial \theta} d\theta dV + \frac{1}{\Delta t} \int_V \delta\theta \rho \frac{\partial U}{\partial \mathbf{g}} \cdot d\mathbf{g} dV \\ - \int_V \delta\mathbf{g} \cdot \frac{\partial \mathbf{f}}{\partial \theta} d\theta dV - \int_V \delta\mathbf{g} \cdot \frac{\partial \mathbf{f}}{\partial \mathbf{g}} \cdot d\mathbf{g} dV \\ - \int_V \delta\theta \frac{\partial r}{\partial \theta} d\theta dV - \int_S \delta\theta \frac{\partial q}{\partial \theta} d\theta dS. \end{aligned}$$

The thermal constitutive behavior for this example is now defined. We assume a constant specific heat for the material. The heat conduction in the material is assumed to be governed by Fourier's law.

The internal thermal energy per unit mass is defined as

$$U = U(\theta),$$

with

$$\frac{\partial U}{\partial \theta} = c,$$

where  $c$  is the specific heat of the material and

$$\frac{\partial U}{\partial \mathbf{g}} = 0.$$

Fourier's law for heat conduction is given as

$$\mathbf{f} = -\mathbf{k} \cdot \mathbf{g},$$

where  $\mathbf{k}$  is the thermal conductivity matrix and  $\mathbf{x}$  is position, so that

$$\frac{\partial \mathbf{f}}{\partial \mathbf{g}} = -\mathbf{k}$$

and

$$\frac{\partial \mathbf{f}}{\partial \theta} = -\frac{\partial \mathbf{k}}{\partial \theta} \cdot \mathbf{g}.$$

The assumption of conductivity without any temperature dependence implies that

$$\frac{\partial \mathbf{f}}{\partial \theta} = 0.$$

No state variables are needed for this material, so the allocation of space for them is not necessary. A thermal user material definition can be used to read in the two constants for our simple case, namely the specific heat,  $c$ , and the coefficient of thermal conductivity,  $k$ , so that

$$\text{PROPS}(1) = k,$$

$$\text{PROPS}(2) = c.$$

```

SUBROUTINE UMATHT(U,DUDT,DUDG,FLUX,DFDT,DFDG,
1 STATEV,TEMP,DTEMP,DTEMDX,TIME,DTIME,PREDEF,DPRED,
2 CMNAME,NTGRD,NSTATV,PROPS,NPROPS,COORDS,PNEWDT,
3 NOEL,NPT,LAYER,KSPT,KSTEP,KINC)
C
      INCLUDE 'ABA_PARAM.INC'
C
      CHARACTER*80 CMNAME
      DIMENSION DUDG(NTGRD),FLUX(NTGRD),DFDT(NTGRD),
1 DFDG(NTGRD,NTGRD),STATEV(NSTATV),DTEMDX(NTGRD),
2 TIME(2),PREDEF(1),DPRED(1),PROPS(NPROPS),COORDS(3)
C
      COND = PROPS(1)
      SPECHT = PROPS(2)
C
      DUDT = SPECHT
      DU = DUDT*DTEMP
      U = U+DU
C
      DO I=1, NTGRD
          FLUX(I) = -COND*DTEMDX(I)
          DFDG(I,I) = -COND
      END DO
C
      RETURN
END

```

### 1.1.39 UMESHMOTION: User subroutine to specify mesh motion constraints during adaptive meshing.

**Product:** Abaqus/Standard

#### References

---

- “Defining ALE adaptive mesh domains in Abaqus/Standard,” Section 12.2.6 of the Abaqus Analysis User’s Manual
- \*ADAPTIVE MESH
- \*ADAPTIVE MESH CONSTRAINT

#### Overview

---

User subroutine **UMESHMOTION**:

- is called at the end of any increment where adaptive meshing is performed (as specified by the FREQUENCY parameter on the \*ADAPTIVE MESH option);
- can be used to define the motion of nodes in an adaptive mesh constraint node set; and
- can call utility routines **GETVRN**, **GETNODETOELEMCONN**, and **GETVRMAVGATNODE** to access results data at the node.

#### Accessing node point data

---

You are provided with access to the values of the node point quantities at the end of the increment through the utility routine **GETVRN** described in “Obtaining node point information,” Section 2.1.9. You can also access values of material point quantities extrapolated to, and averaged, at nodes at the end of the increment through the utility routine **GETVRMAVGATNODE** described in “Obtaining material point information averaged at a node,” Section 2.1.8. **GETVRMAVGATNODE** requires the list of elements attached to the node, which is obtained by calling the utility routine **GETNODETOELEMCONN** described in “Obtaining node to element connectivity,” Section 2.1.10.

#### User subroutine interface

---

```

SUBROUTINE UMESHMOTION(UREF,ULOCAL,NODE,NNDOF,
*      LNODETYPE,ALOCAL,NDIM,TIME,DTIME,PNEWDT,
*      KSTEP,KINC,KMESHSWEEP,JMATYP,JGVBLOCK,LSMOOTH)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION ULOCAL(NDIM),JELEMLIST(*)
DIMENSION ALOCAL(NDIM,*),TIME(2)

```

## UMESHMOTION

```
DIMENSION JMATYP(*),JGVBLOCK(*)  
C
```

*user coding to define **ULOCAL**  
and, optionally **PNEWDT***

```
RETURN  
END
```

### Variable to be defined

---

#### **ULOCAL**

Components of the mesh displacement or velocity of the adaptive mesh constraint node, described in the coordinate system **ALOCAL**. **ULOCAL** will be passed into the routine as values determined by the mesh smoothing algorithm. All components of the mesh displacement or velocity will be applied; i.e., you do not have the ability to select the directions in which the mesh displacement should be applied.

### Variables that can be updated

---

#### **PNEWDT**

Ratio of suggested new time increment to the time increment currently being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **UMESHMOTION**.

The suggested new time increment provided to the automatic time integration algorithms is  $\text{PNEWDT} \times \text{DTIME}$ , where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this increment.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** greater than 1.0 will be ignored and values of **PNEWDT** less than 1.0 will cause the job to terminate.

#### **LSMOOTH**

Flag specifying that surface smoothing be applied after application of the mesh motion constraint. Set **LSMOOTH** to 1 to enable surface smoothing. When this flag is set, the constraint defined in **ULOCAL** will be modified by the smoothing algorithm. In cases where **ULOCAL** describes mesh motion normal to a surface, the smoothing will have a minor impact on this normal component of mesh motion.

### Variables passed in for information

---

#### **UREF**

The value of the user-specified displacement or velocity provided as part of the adaptive mesh constraint definition. This value is updated based on any amplitude definitions used with the adaptive mesh constraint or default ramp amplitude variations associated with the current step.

**NODE**

Node number.

**NNDOF**

Number of degrees of freedom at the node.

**LNODETYPE**

Node type flag.

**LNODETYPE**=1 indicates that the node is on the interior of the adaptive mesh region.

**LNODETYPE**=2 indicates that the node is involved in a tied constraint.

**LNODETYPE**=3 indicates that the node is at the corner of the boundary of an adaptive mesh region.

**LNODETYPE**=4 indicates that the node lies on the edge of a boundary of an adaptive mesh region.

**LNODETYPE**=5 indicates that the node lies on a flat surface on a boundary of the adaptive mesh region.

**LNODETYPE**=6 indicates that the node participates in a constraint (other than a tied constraint) as a master node.

**LNODETYPE**=7 indicates that the node participates in a constraint (other than a tied constraint) as a slave node.

**LNODETYPE**=10 indicates that a concentrated load is applied to the node.

**ALOCAL**

Local coordinate system aligned with the tangent to the adaptive mesh domain at the node. If the node is on the interior of the adaptive mesh domain, **ALOCAL** will be set to the identity matrix. In other cases the 1-direction is along an edge or in the plane of a flat surface. When **NDIM**=2, the 2-direction is normal to the surface. When **NDIM**=3, the 2-direction also lies in the plane of a flat surface or is arbitrary if the node is on an edge. When **NDIM**=3 the 3-direction is normal to the surface or is arbitrary if the node is on an edge.

**NDIM**

Number of coordinate dimensions.

**TIME (1)**

Current value of step time.

**TIME (2)**

Current value of total time.

**DTIME**

Time increment.

**KSTEP**

Step number.

**KINC**

Increment number.

## UMESHMOTION

### **KMESHSWEEP**

Mesh sweep number.

### **JMATYP**

Variable that must be passed into the **GETVRMAVGATNODE** utility routine to access local results at the node.

### **JGVBLOCK**

Variable that must be passed into the **GETVRN**, **GETNODETOELEMCONN**, and **GETVRMAVGATNODE** utility routines to access local results at the node.

## 1.1.40 UMOTION: User subroutine to specify motions during cavity radiation heat transfer analysis or steady-state transport analysis.

**Product:** Abaqus/Standard

### References

---

- “Cavity radiation,” Section 37.1.1 of the Abaqus Analysis User’s Manual
- “Steady-state transport analysis,” Section 6.4.1 of the Abaqus Analysis User’s Manual
- \*MOTION
- \*TRANSPORT VELOCITY

### Overview

---

User subroutine **UMOTION**:

- can be used either to define the magnitude of the translational motion for degrees of freedom specified as a predefined field in a cavity radiation heat transfer analysis or to define the magnitude of the rotational velocity in a steady-state transport step; and
- will overwrite any motion or transport velocity magnitudes if they are defined directly (and possibly modified by including an amplitude reference) outside the user subroutine.

### User subroutine interface

---

```

SUBROUTINE UMOTION(U,KSTEP,KINC,TIME,NODE,JDOF)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION U,TIME(2)
C

```

*user coding to define U*

```

RETURN
END

```

---

## Variable to be defined

---

### **U**

Total value of the component of the translation due to prescribed motion for the degree of freedom specified by **JDOF**. **U** will be passed into the routine as the value defined by any magnitude and/or amplitude specification in the motion definition for the degree of freedom **JDOF**. The total value of the translation must be given in user subroutine **UMOTION**, regardless of the type of motion defined (displacement or velocity).

When used in conjunction with a steady-state transport analysis, **U** defines the magnitude of the rotational velocity. In such a case **JDOF** is passed in as **0**.

---

## Variables passed in for information

---

### **KSTEP**

Step number.

### **KINC**

Increment number.

### **TIME (1)**

Current value of step time.

### **TIME (2)**

Current value of total time.

### **NODE**

Node number.

### **JDOF**

Degree of freedom. When used in a steady-state transport analysis, **JDOF** is passed in as **0**.

## 1.1.41 UMULLINS: User subroutine to define damage variable for the Mullins effect material model.

**Product:** Abaqus/Standard

### References

---

- “Mullins effect,” Section 19.6.1 of the Abaqus Analysis User’s Manual
- \*MULLINS EFFECT
- “Mullins effect and permanent set,” Section 2.2.3 of the Abaqus Verification Manual

### Overview

---

User subroutine **UMULLINS**:

- can be used to define the damage variable for the Mullins effect material model, including the use of the Mullins effect approach to model energy dissipation in elastomeric foams;
- will be called at all material calculation points of elements for which the material definition contains a user-defined Mullins effect; and
- should be used when you do not want to use the form of the damage variable,  $\eta$ , that is used by Abaqus/Standard.

### User subroutine interface

---

```

SUBROUTINE UMULLINS (NUMPROPS , PROPS , UMAXNEW , UMAXOLD , SEDDEV ,
1 ETA , DETADW , DMGDISSOLD , DMGDISSNEW , SENERNEW , NUMSTATEV , STATEV ,
2 TEMP , DTEMP , NUMFIELDV , FIELDV , FIELDVINC , CMNAME , LINPER)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
DIMENSION PROPS(*) , STATEV(*) , FIELDV(*) , FIELDVINC(*)

user coding to define ETA, DETADW,
and, optionally, DMGDISSNEW, SENERNEW, STATEV

RETURN
END

```

---

**Variables to be defined**

---

**ETA**

The damage variable,  $\eta$ .

**DETADW**

The derivative of the damage variable with respect to the elastic strain energy density of the undamaged material,  $\frac{d\eta}{dU}$ . This quantity is needed for the Jacobian of the overall system of equations and needs to be defined accurately to ensure good convergence characteristics.

---

**Variables that can be updated**

---

**DMGDISSNEW**

The energy dissipation density at the end of the increment. This quantity can be defined either in total form or in an incremental manner using the old value of the damage dissipation **DMGDISSOLD** and the increment in damage dissipation. This quantity is used for output purposes only.

**SENERNEW**

The recoverable strain energy density at the end of the increment. This quantity is used for output purposes only.

**STATEV**

Array containing the user-defined solution-dependent state variables at this point. These are supplied as values at the start of the increment or as values updated by other user subroutines (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual) and must be returned as values at the end of the increment.

---

**Variables passed in for information**

---

**UMAXNEW**

The value, at the end of the increment, of the maximum primary strain energy density over its entire deformation history.

**UMAXOLD**

The value, at the beginning of the increment, of the maximum primary strain energy density over its entire deformation history.

**SEDDEV**

The value, at the end of the increment, of the deviatoric primary strain energy density when the primary material behavior is hyperelastic. The value, at the end of the increment, of the total primary strain energy density when the primary material behavior is hyperfoam.

**DMGDISSOLD**

The value of energy dissipated at the beginning of the increment.

**CMNAME**

User-specified material name, left justified.

**NUMSTATEV**

Number of solution-dependent state variables associated with this material (defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NUMPROPS**

Number of material properties entered for this user-defined hyperelastic material.

**PROPS**

Array of material properties entered for this user-defined hyperelastic material.

**TEMP**

Temperature at the start of the increment.

**DTEMP**

Increment of temperature.

**NUMFIELDV**

Number of field variables.

**FIELDV**

Array of interpolated values of predefined field variables at this material point at the beginning of the increment based on the values read in at the nodes (initial values at the beginning of the analysis and current values during the analysis).

**FIELDVINC**

Array of increments of predefined field variables at this material point for this increment; this includes any values updated by user subroutine **USDFLD**.

**LINPER**

Linear perturbation flag. **LINPER**=1 if the step is a linear perturbation step. **LINPER**=0 if the step is a general step.



## 1.1.42 UPOREP: User subroutine to define initial fluid pore pressure.

**Product:** Abaqus/Standard

### References

---

- “Initial conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.2.1 of the Abaqus Analysis User’s Manual
- “Coupled pore fluid diffusion and stress analysis,” Section 6.8.1 of the Abaqus Analysis User’s Manual
- \*INITIAL CONDITIONS

### Overview

---

User subroutine **UPOREP**:

- allows for the specification of the initial pore pressure values of a porous medium;
- can be used to define initial pore pressure values as functions of nodal coordinates and/or node numbers; and
- will be called to define initial fluid pore pressure values at all nodes of a coupled pore fluid diffusion and stress analysis whenever user-defined initial pore pressure conditions are specified.

### User subroutine interface

---

```
SUBROUTINE UPOREP (UW0 , COORDS , NODE)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION COORDS (3)
C
```

*user coding to define UW0*

```
RETURN
END
```

### Variable to be defined

---

#### UW0

Initial fluid pore pressure.

## **Variables passed in for information**

---

### **COORDS**

An array containing the current coordinates of this node.

### **NODE**

Node number.

## 1.1.43 UPRESS: User subroutine to specify prescribed equivalent pressure stress conditions.

**Product:** Abaqus/Standard

### References

---

- “Mass diffusion analysis,” Section 6.9.1 of the Abaqus Analysis User’s Manual
- \*PRESSURE STRESS
- “**UTEMP**, **UFIELD**, **UMASF**, and **UPRESS**,” Section 4.1.25 of the Abaqus Verification Manual

### Overview

---

User subroutine **UPRESS**:

- allows you to prescribe equivalent pressure stress values at the nodes of a model;
- will be called in a mass diffusion analysis whenever a current value of equivalent pressure stress is needed for a node that has user-defined pressure stress conditions;
- can be used to modify any pressure stresses read in from a results file; and
- ignores any equivalent pressure stresses provided for the associated pressure stress definition outside the user subroutine.

### User subroutine interface

---

```
SUBROUTINE UPRESS(PRESS,KSTEP,KINC,TIME,NODE,COORDS)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION TIME(2), COORDS(3)
C
```

*user coding to define PRESS*

```
RETURN
END
```

### Variable to be defined

---

#### PRESS

Total value of the equivalent pressure stress at the node.

You may have also requested equivalent pressure stress to be set in one of two other ways: from a previously generated results file or via direct data input. When **PRESS** is passed into user subroutine **UPRESS**, it will contain equivalent pressure stresses obtained from the results file only. You can modify these values within this routine. Any values given as direct data input will be ignored.

### **Variables passed in for information**

---

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Current value of step time.

**TIME (2)**

Current value of total time.

**NODE**

Node number.

**COORDS**

An array containing the coordinates of this node.

### 1.1.44 UPSD: User subroutine to define the frequency dependence for random response loading.

**Product:** Abaqus/Standard

#### References

---

- “Random response analysis,” Section 6.3.11 of the Abaqus Analysis User’s Manual
- \*RANDOM RESPONSE
- \*PSD-DEFINITION
- “Random response to jet noise excitation,” Section 1.4.10 of the Abaqus Benchmarks Manual

#### Overview

---

User subroutine **UPSD**:

- will be called once for each frequency at which calculations will be made during a random response analysis if the frequency function is defined in a user subroutine;
- is used to define complicated frequency dependencies for the cross-spectral density matrix of the random loading; and
- ignores any data given for the associated frequency function outside the user subroutine.

#### User subroutine interface

---

```

SUBROUTINE UPSD (PSD, PSDR, PSDI, FREQ, KSTEP)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 PSD

```

*user coding to define PSDR and PSDI*

```

RETURN
END

```

#### Variables to be defined

---

##### PSDR

Real part of the frequency function at this frequency.

**PSDI**

Imaginary part of the frequency function at this frequency.

**Variables passed in for information**

---

**PSD**

User-specified name for this frequency function definition, left justified.

**FREQ**

Frequency, in radians per time.

**KSTEP**

Step number.

## 1.1.45 URDFIL: User subroutine to read the results file.

**Product:** Abaqus/Standard

### References

---

- “Results file output format,” Section 5.1.2 of the Abaqus Analysis User’s Manual
- “Accessing the results file information,” Section 5.1.3 of the Abaqus Analysis User’s Manual
- “Utility routines for accessing the results file,” Section 5.1.4 of the Abaqus Analysis User’s Manual

### Overview

---

User subroutine **URDFIL**:

- can be used to access the results file during an analysis;
- is called at the end of any increment in which new information is written to the results file;
- must call the utility routine **DBFILE** to read records from the results file (see “Utility routines for accessing the results file,” Section 5.1.4 of the Abaqus Analysis User’s Manual);
- can call the utility routine **POSFILE** to read from the results file starting at a specified step and increment as opposed to the beginning of the file, which would otherwise be done (see “Utility routines for accessing the results file,” Section 5.1.4 of the Abaqus Analysis User’s Manual);
- can force an analysis to terminate upon completion of a call by means of the variable **LSTOP**;
- allows the last increment written to the results file to be overwritten by means of the variable **LOVRWRT**; and
- allows access to the complete results file in a restarted job if the new results file is being appended to the old results file (see the description of the execution option **fil** in “Abaqus/Standard, Abaqus/Explicit, and Abaqus/CFD execution,” Section 3.2.2 of the Abaqus Analysis User’s Manual).

### User subroutine interface

---

```

SUBROUTINE URDFIL(LSTOP,LOVRWRT,KSTEP,KINC,DTIME,TIME)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION ARRAY(513),JRRAY(NPRECD,513),TIME(2)
EQUIVALENCE (ARRAY(1),JRRAY(1,1))

```

*user coding to read the results file*

```

RETURN
END

```

---

## Variables to be defined

### In all cases

#### **LSTOP**

Flag to indicate whether an analysis should continue. The analysis will be terminated if **LSTOP** is set to 1. Otherwise, the analysis will continue.

#### **LOVRWRT**

Flag to indicate that the information written to the results file for the increment can be overwritten. If **LOVRWRT** is set to 1, information for the current increment will be overwritten by information written to the results file in a subsequent increment unless the current increment is the final increment written to the results file. The purpose of this flag is to reduce the size of the results file by allowing information for an increment to be overwritten by information for a subsequent increment.

#### **DTIME**

Time increment. This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus (if automatic time incrementation is chosen). It is passed in as the value of the next time increment to be taken and can be updated to increase or reduce the time increment. If automatic time incrementation is not selected in the analysis procedure, updated values of **DTIME** are ignored.

### Only if utility routine **POSFIL** is called

#### **NSTEP**

Desired step at which file reading will begin via utility routine **DBFILE**. If **NSTEP** is set to 0, the first available step will be read.

#### **NINC**

Desired increment at which file reading will begin via utility routine **DBFILE**. If **NINC** is set to 0, the first available increment of the specified step will be read.

---

## Variables passed in for information

#### **KSTEP**

Step number.

#### **KINC**

Increment number.

#### **TIME (1)**

Value of the step time at the end of the increment.

#### **TIME (2)**

Value of the total time at the end of the increment.

---

**Example: Terminating an analysis upon exceeding a Mises stress limit**

---

The example below reads the values of Mises stress for the current increment from record 12 in the results file and terminates the analysis if any of the values of Mises stress written to the results file exceed  $2.09 \times 10^8$ . Here, **POSFIL** is used to position you to read from the current increment.

```

SUBROUTINE URDFIL(LSTOP,LOVRWRT,KSTEP,KINC,DTIME,TIME)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION ARRAY(513),JRRAY(NPRECD,513),TIME(2)
EQUIVALENCE (ARRAY(1),JRRAY(1,1))
PARAMETER(TOL=2.09D8)
C
C FIND CURRENT INCREMENT.
C
CALL POSFIL(KSTEP,KINC,ARRAY,JRCD)
DO K1=1,999999
    CALL DBFILE(0,ARRAY,JRCD)
    IF (JRCD .NE. 0) GO TO 110
    KEY=JRRAY(1,2)
C
C RECORD 12 CONTAINS VALUES FOR SINV
C
    IF (KEY.EQ.12) THEN
        IF (ARRAY(3).GT.TOL) THEN
            LSTOP=1
            GO TO 110
        END IF
    END IF
END DO
110 CONTINUE
C
RETURN
END

```

---

**Example: Terminating an analysis when the maximum Mises stress value stops increasing**

---

This example demonstrates the use of **URDFIL** and **POSFIL** to stop an analysis when the maximum value of Mises stress in the model does not increase from one increment in the results file to the next. A data statement is used to save the maximum Mises stress value from the last increment. **LOVRWRT** is also used in this case to overwrite an increment in the results file once it has been read in **URDFIL**.

The subroutine shown below must be modified to define the maximum Mises stress in the data statement each time a job is restarted. This can be avoided by removing the **LOVRWRT=1** statement and recoding the routine to read both the previous and the current increment to check that the Mises stress increases from one increment to the next (in this case you must correctly handle the first increment written to the results file as there will be no previous increment). The results file must also be properly appended on restart if you wish to compare the values of Mises stress between the first increment of a restart and the final increment of the job being restarted. This approach has the disadvantage that the results file may become quite large, as no information in the file will be overwritten.

```

SUBROUTINE URDFIL(LSTOP,LOVRWRT,KSTEP,KINC,DTIME,TIME)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION ARRAY(513),JRRAY(NPRECD,513),TIME(2)
EQUIVALENCE (ARRAY(1),JRRAY(1,1))
C
C INITIALIZE THE OLD MAXIMUM. FOR A JOB THAT IS BEING RESTARTED
C THIS VALUE SHOULD BE SET TO THE MAXIMUM MISES STRESS IN THE
C ORIGINAL ANALYSIS.
C
DATA OLDMAX/-1.D0/
C
CURRMAX = 0.D0
C
C FIND CURRENT INCREMENT.
C
CALL POSFIL(KSTEP,KINC,ARRAY,JRCD)
C
C SEARCH FOR THE HIGHEST VALUE OF MISES STRESS
C AND STORE THIS IN CURRMAX
C
DO K1=1,999999
    CALL DBFILE(0,ARRAY,JRCD)
    IF (JRCD.NE.0) GO TO 110
    KEY=JRRAY(1,2)
    IF (KEY.EQ.12) THEN
        IF (ARRAY(3).GT.CURRMAX) CURRMAX=ARRAY(3)
    END IF
END DO
110 CONTINUE
C
C COMPLETED READING OF CURRENT INCREMENT. NOW CHECK TO
C SEE IF VALUE OF MISES STRESS HAS INCREASED SINCE

```

```
C LAST INCREMENT
C
IF (CURRMAX.LE.OLDMAX) LSTOP=1
OLDMAX=CURRMAX
LOVRWRT=1
C
RETURN
END
```



## 1.1.46 USDFLD: User subroutine to redefine field variables at a material point.

**Product:** Abaqus/Standard

### References

---

- “Obtaining material point information in an Abaqus/Standard analysis,” Section 2.1.6
- “Material data definition,” Section 18.1.2 of the Abaqus Analysis User’s Manual
- \*USER DEFINED FIELD
- “Damage and failure of a laminated composite plate,” Section 1.1.14 of the Abaqus Example Problems Manual
- “**USDFLD**,” Section 4.1.24 of the Abaqus Verification Manual

### Overview

---

User subroutine **USDFLD**:

- allows you to define field variables at a material point as functions of time or of any of the available material point quantities listed in the Output Variable Identifiers table (“Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual) except the user-defined output variables UVARM and UVARM $n$ ;
- can be used to introduce solution-dependent material properties since such properties can easily be defined as functions of field variables;
- will be called at all material points of elements for which the material definition includes user-defined field variables;
- must call utility routine **GETVRM** to access material point data;
- can use and update state variables; and
- can be used in conjunction with user subroutine **UFIELD** to prescribe predefined field variables.

### Explicit solution dependence

---

Since this routine provides access to material point quantities only at the start of the increment, the solution dependence introduced in this way is explicit: the material properties for a given increment are not influenced by the results obtained during the increment. Hence, the accuracy of the results depends on the size of the time increment. Therefore, you can control the time increment in this routine by means of the variable **PNEWDT**.

### Defining field variables

---

Before user subroutine **USDFLD** is called, the values of the field variables at the material point are calculated by interpolation from the values defined at the nodes. Any changes to the field variables in the user subroutine are local to the material point: the nodal field variables retain the values defined

as initial conditions, predefined field variables, or in user subroutine **UFIELD**. The values of the field variables defined in this routine are used to calculate values of material properties that are defined to depend on field variables and are passed into other user subroutines that are called at the material point, such as the following:

- **CREEP**
- **HETVAL**
- **UEXPAN**
- **UHARD**
- **UHYPEL**
- **UMAT**
- **UMATHT**
- **UTRS**

Output of the user-defined field variables at the material points can be obtained with the element integration point output variable FV (see “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual).

## Accessing material point data

---

You are provided with access to the values of the material point quantities at the start of the increment (or in the base state in a linear perturbation step) through the utility routine **GETVRM** described in “Obtaining material point information in an Abaqus/Standard analysis,” Section 2.1.6. The values of the material point quantities are obtained by calling **GETVRM** with the appropriate output variable keys. The values of the material point data are recovered in the arrays **ARRAY**, **JARRAY**, and **FLGRAY** for floating point, integer, and character data, respectively. You may not get values of some material point quantities that have not been defined at the start of the increment; e.g., ER.

## State variables

---

Since the redefinition of field variables in **USDFLD** is local to the current increment (field variables are restored to the values interpolated from the nodal values at the start of each increment), any history dependence required to update material properties by using this subroutine must be introduced with user-defined state variables.

The state variables can be updated in **USDFLD** and then passed into other user subroutines that can be called at this material point, such as those listed above. You specify the number of such state variables, as shown in the example at the end of this section (see also “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

## User subroutine interface

---

```
SUBROUTINE USDFLD (FIELD , STATEV , PNEWDT , DIRECT , T , CELENT ,
1 TIME , DTIME , CMNAME , ORNAME , NFIELD , NSTATV , NOEL , NPT , LAYER ,
```

```

2 KSPT,KSTEP,KINC,NDI,NSHR,COORD,MAC,JMATYP,MATLAYO,LACCFLA)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME,ORNAME
CHARACTER*3 FLGRAY(15)
DIMENSION FIELD(NFIELD),STATEV(NSTATV),DIRECT(3,3),
1 T(3,3),TIME(2)
DIMENSION ARRAY(15),JARRAY(15),MAC(*),JMATYP(*),COORD(*)

```

*user coding to define **FIELD** and, if necessary, **STATEV** and **PNEWDT***

```

RETURN
END

```

## Variable to be defined

---

### **FIELD (NFIELD)**

An array containing the field variables at the current material point. These are passed in with the values interpolated from the nodes at the end of the current increment, as specified with initial condition definitions, predefined field variable definitions, or user subroutine **UFIELD**. The interpolation is performed using the same scheme used to interpolate temperatures: an average value is used for linear elements; an approximate linear variation is used for quadratic elements (also see “Solid (continuum) elements,” Section 25.1.1 of the Abaqus Analysis User’s Manual). The updated values are used to calculate the values of material properties that are defined to depend on field variables and are passed into other user subroutines (**CREEP**, **HETVAL**, **UEXPAN**, **UHARD**, **UHYPER**, **UMAT**, **UMATHT**, and **UTRS**) that are called at this material point.

## Variables that can be updated

---

### **STATEV (NSTATV)**

An array containing the solution-dependent state variables. These are passed in as the values at the beginning of the increment. In all cases **STATEV** can be updated in this subroutine, and the updated values are passed into other user subroutines (**CREEP**, **HETVAL**, **UEXPAN**, **UMAT**, **UMATHT**, and **UTRS**) that are called at this material point. The number of state variables associated with this material point is defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

### **PNEWDT**

Ratio of suggested new time increment to the time increment being used (**DTIME**, see below). This variable allows you to provide input to the automatic time incrementation algorithms in Abaqus/Standard (if automatic time incrementation is chosen).

**PNEWDT** is set to a large value before each call to **USDFLD**.

If **PNEWDT** is redefined to be less than 1.0, Abaqus/Standard must abandon the time increment and attempt it again with a smaller time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines that allow redefinition of **PNEWDT** for this iteration.

If **PNEWDT** is given a value that is greater than 1.0 for all calls to user subroutines for this iteration and the increment converges in this iteration, Abaqus/Standard may increase the time increment. The suggested new time increment provided to the automatic time integration algorithms is **PNEWDT** × **DTIME**, where the **PNEWDT** used is the minimum value for all calls to user subroutines for this iteration.

If automatic time incrementation is not selected in the analysis procedure, values of **PNEWDT** that are greater than 1.0 will be ignored and values of **PNEWDT** that are less than 1.0 will cause the job to terminate.

## Variables passed in for information

---

### **DIRECT(3,3)**

An array containing the direction cosines of the material directions in terms of the global basis directions. **DIRECT(1,1)**, **DIRECT(2,1)**, **DIRECT(3,1)** give the (1, 2, 3) components of the first material direction; **DIRECT(1,2)**, **DIRECT(2,2)**, **DIRECT(3,2)** give the second material direction, etc. For shell and membrane elements, the first two directions are in the plane of the element and the third direction is the normal. This information is not available for beam elements.

### **T(3,3)**

An array containing the direction cosines of the material orientation components relative to the element basis directions. This is the orientation that defines the material directions (**DIRECT**) in terms of the element basis directions. For continuum elements **T** and **DIRECT** are identical. For shell and membrane elements **T(1,1)** =  $\cos \theta$ , **T(1,2)** =  $-\sin \theta$ , **T(2,1)** =  $\sin \theta$ , **T(2,2)** =  $\cos \theta$ , **T(3,3)** = 1.0, and all other components are zero, where  $\theta$  is the counterclockwise rotation around the normal vector that defines the orientation. If no orientation is used, **T** is an identity matrix. Orientation is not available for beam elements.

### **CELENT**

Characteristic element length. This is a typical length of a line across an element for a first-order element; it is half of the same typical length for a second-order element. For beams and trusses it is a characteristic length along the element axis. For membranes and shells it is a characteristic length in the reference surface. For axisymmetric elements it is a characteristic length in the  $(r, z)$  plane only.

### **TIME(1)**

Value of step time at the beginning of the current increment.

### **TIME(2)**

Value of total time at the beginning of the current increment.

**DTIME**

Time increment.

**CMNAME**

User-specified material name, left justified.

**ORNAME**

User-specified local orientation name, left justified.

**NFIELD**

Number of field variables defined at this material point.

**NSTATV**

User-defined number of solution-dependent state variables (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**KSTEP**

Step number.

**KINC**

Increment number.

**NDI**

Number of direct stress components at this point.

**NSHR**

Number of shear stress components at this point.

**COORD**

Coordinates at this material point.

**JMAC**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**JMATYP**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**MATLAYO**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**LACCFLA**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**Example: Damaged elasticity model**

---

Included below is an example of user subroutine **USDFLD**. In this example a truss element is loaded in tension. A damaged elasticity model is introduced: the modulus decreases as a function of the maximum tensile strain that occurred during the loading history. The maximum tensile strain is stored as a solution-dependent state variable—see “Defining solution-dependent field variables” in “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual.

**Input file**

```
*HEADING
  DAMAGED ELASTICITY MODEL WITH USER SUBROUTINE USDFLD
*ELEMENT, TYPE=T2D2, ELSET=ONE
  1, 1, 2
*NODE
  1, 0., 0.
  2, 10., 0.
*SOLID SECTION, ELSET=ONE, MATERIAL=ELASTIC
  1.
*MATERIAL, NAME=ELASTIC
*ELASTIC, DEPENDENCIES=1
** Table of modulus values decreasing as a function
** of field variable 1.
  2000., 0.3, 0., 0.00
  1500., 0.3, 0., 0.01
  1200., 0.3, 0., 0.02
  1000., 0.3, 0., 0.04
*USER DEFINED FIELD
*DEPVAR
  1
*BOUNDARY
  1, 1, 2
  2, 2
*STEP
*STATIC
```

```

0.1, 1.0, 0.0, 0.1
*CLOAD
2, 1, 20.
*END STEP
*STEP
*STATIC
0.1, 1.0, 0.0, 0.1
*CLOAD
2, 1, 0.
*END STEP
*STEP, INC=20
*STATIC
0.1, 2.0, 0.0, 0.1
*CLOAD
2, 1, 40.
*END STEP

```

**User subroutine**

```

SUBROUTINE USDFLD(FIELD,STATEV,PNEWDT,DIRECT,T,CELENT,
1 TIME,DTIME,CMNAME,ORNAME,NFIELD,NSTATV,NOEL,NPT,LAYER,
2 KSPT,KSTEP,KINC,NDI,NSHR,COORD,JMAC,JMATYP,MATLayo,
3 LACCFLA)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME,ORNAME
CHARACTER*3 FLGRAY(15)
DIMENSION FIELD(NFIELD),STATEV(NSTATV),DIRECT(3,3),
1 T(3,3),TIME(2)
DIMENSION ARRAY(15),JARRAY(15),JMAC(*),JMATYP(*),
1 COORD(*)
C
C Absolute value of current strain:
CALL GETVRM('E',ARRAY,JARRAY,FLGRAY,JRCD,JMAC,JMATYP,
MATLayo,LACCFLA)
EPS = ABS( ARRAY(1) )
C Maximum value of strain up to this point in time:
CALL GETVRM('SDV',ARRAY,JARRAY,FLGRAY,JRCD,JMAC,JMATYP,
MATLayo,LACCFLA)
EPSMAX = ARRAY(1)
C Use the maximum strain as a field variable
FIELD(1) = MAX( EPS , EPSMAX )

```

```
C Store the maximum strain as a solution dependent state
C variable
      STATEV(1) = FIELD(1)
C If error, write comment to .DAT file:
      IF(JRCD.NE.0)THEN
          WRITE(6,*) 'REQUEST ERROR IN USDFLD FOR ELEMENT NUMBER ',
1          NOEL,'INTEGRATION POINT NUMBER ',NPT
      ENDIF
C
      RETURN
END
```

### 1.1.47 UTEMP: User subroutine to specify prescribed temperatures.

**Product:** Abaqus/Standard

#### References

---

- “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual
- \*TEMPERATURE
- “LE11: Solid cylinder/taper/sphere—temperature loading,” Section 4.2.11 of the Abaqus Benchmarks Manual
- “**UTEMP**, **UFIELD**, **UMASFL**, and **UPRESS**,” Section 4.1.25 of the Abaqus Verification Manual

#### Overview

---

User subroutine **UTEMP**:

- allows you to prescribe temperatures at the nodes of a model;
- will be called whenever a current value of temperature is needed for a node that is listed under a user-defined temperature field definition;
- ignores any temperatures provided for the associated temperature field definition outside the user subroutine; and
- can be used to modify any temperatures read in from a results file.

#### User subroutine interface

---

```

SUBROUTINE UTEMP (TEMP ,NSECPT ,KSTEP ,KINC ,TIME ,NODE ,COORDS)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION TEMP (NSECPT) , TIME (2) , COORDS (3)
C

```

*user coding to define TEMP*

```

RETURN
END

```

**Variable to be defined**

---

**TEMP (NSECPT)**

Array of temperature values at node number **NODE**. If the node is not connected to a beam or shell element, only one value of temperature must be returned (**NSECPT**=1). Otherwise, the number of temperatures to be returned depends on the mode of temperature and field variable input selected for the beam or shell section. The following cases are possible:

1. Temperatures and field variables for a beam section are given as values at the points shown in the beam section descriptions. The number of values required, **NSECPT**, is determined by the particular section type specified, as described in “Beam cross-section library,” Section 26.3.9 of the Abaqus Analysis User’s Manual.
2. Temperatures and field variables are given as values at  $n$  equally spaced points through each layer of a shell section. The number of values required, **NSECPT**, is equal to  $n$ .
3. Temperatures and field variables for a beam section are given as values at the origin of the cross-section together with gradients with respect to the 2-direction and, for three-dimensional beams, the 1-direction of the section; or temperatures and field variables for a shell section are given as values at the reference surface together with gradients with respect to the thickness. The number of values required, **NSECPT**, is 3 for three-dimensional beams, 2 for two-dimensional beams, and 2 for shells. Give the midsurface value first, followed by the first and (if necessary) second gradients, as described in “Beam elements,” Section 26.3 of the Abaqus Analysis User’s Manual, and “Shell elements,” Section 26.6 of the Abaqus Analysis User’s Manual.

You can also request temperatures to be set in one of two other ways: from a previously generated results file or via direct data input. When array **TEMP** is passed into user subroutine **UTEMP**, it will contain temperatures obtained from the results file only. You can modify these values within this routine. Any values given as direct data input will be ignored.

**Variables passed in for information**

---

**NSECPT**

Maximum number of section values required for any node in the model.

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Current value of step time.

**TIME (2)**

Current value of total time.

**NODE**

Node number.

**COORDS**

An array containing the current coordinates of this point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the node.



## 1.1.48 UTRACLOAD: User subroutine to specify nonuniform traction loads.

**Product:** Abaqus/Standard

### References

---

- “Distributed loads,” Section 30.4.3 of the Abaqus Analysis User’s Manual
- \*DLOAD
- \*DSLOAD
- “Distributed traction and edge loads,” Section 1.4.17 of the Abaqus Verification Manual

### Overview

---

User subroutine **UTRACLOAD**:

- can be used to define the variation of the distributed traction load magnitude as a function of position, time, element number, load integration point number, etc.;
- if needed, can be used to define the *initial* loading direction for the distributed traction load as a function of position, element number, load integration point number, etc.;
- will be called at each load integration point for each element-based, edge-based, or surface-based nonuniform distributed traction load definition during stress analysis;
- cannot be used in mode-based procedures to describe the time variation of the load; and
- ignores any amplitude references that may appear with the associated step definition or nonuniform distributed traction load definition.

### User subroutine interface

---

```

SUBROUTINE UTRACLOAD (ALPHA,T_USER,KSTEP,KINC,TIME,NOEL,NPT,
1 COORDS,DIRCOS,JLTYP,SNAME)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION T_USER(3), TIME(2), COORDS(3), DIRCOS(3,3)
CHARACTER*80 SNAME

```

*user coding to define ALPHA and T\_USER*

```

RETURN
END

```

---

## Variables to be defined

**ALPHA**

Magnitude of the distributed traction load. Units are  $\text{FL}^{-2}$  for surface loads,  $\text{FL}^{-1}$  for edge loads, and F for edge moments. **ALPHA** is passed into the routine as the magnitude of the load specified as part of the element-based or surface-based distributed load definition. If the magnitude is not defined, **ALPHA** is passed in as zero. For a static analysis that uses the modified Riks method (“Unstable collapse and postbuckling analysis,” Section 6.2.4 of the Abaqus Analysis User’s Manual) **ALPHA** must be defined as a function of the load proportionality factor,  $\lambda$ . The distributed load magnitude is not available for output purposes.

**T\_USER**

Loading direction of the distributed traction load. **T\_USER** is passed into the routine as the load direction specified as part of the element-based or surface-based distributed load definition. The vector **T\_USER** passed out of the subroutine is used as the initial loading direction  $t_{user}$  discussed in “Distributed loads,” Section 30.4.3 of the Abaqus Analysis User’s Manual. The direction of **T\_USER** as defined by the subroutine should not change during a step. If it does, convergence difficulties might arise. Load directions are needed only for a nonuniform general surface traction, shear surface traction, and general edge traction. If a direction is defined for the nonuniform normal edge traction, shear edge traction, transverse edge traction, or edge moment, it will be ignored. See “Distributed loads,” Section 30.4.3 of the Abaqus Analysis User’s Manual, for details.

---

## Variables passed in for information

**KSTEP**

Step number.

**KINC**

Increment number.

**TIME (1)**

Current value of step time or current value of the load proportionality factor,  $\lambda$ , in a Riks step.

**TIME (2)**

Current value of total time.

**NOEL**

User-defined element number.

**NPT**

Load integration point number within the element or on the element’s surface, depending on the load type.

**COORDS**

An array containing the coordinates of the load integration point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

**DIRCOS**

Orientation of the face or edge in the reference configuration. For three-dimensional facets the first and second columns are the normalized local directions in the plane of the surface, and the third column is the normal to the face. For solid elements the normal points inward, which is the negative of what is defined in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual; for shell elements the normal definition is consistent with the convention. For two-dimensional facets the first column is the normalized tangent, the second column is the facet normal, and the third column is not used. For three-dimensional shell edges the first column is the tangent to the shell edge (shear direction), the second column is the in-plane normal (normal direction), and the third column is the normal to the plane of the shell (transverse direction).

**JLTYP**

Identifies the load type for which this call to **UTRACLOAD** is being made. The load type may be an element-based surface load, an edge-based load, or a surface-based load. This variable identifies the element face or edge for which this call to **UTRACLOAD** is being made. This information is useful when several different nonuniform distributed loads are being imposed on an element at the same time. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for element face and edge identification. The load labels are shown in Table 1.1.48–1. For surface- or edge-based loading (TRSHRNU, TRVECNU, EDLDNU, EDNORNU, EDSHRNU, EDTRANU, EDMOMNU),  $j$  in the load type identifies the face or edge of the element underlying the surface.

**Table 1.1.48–1 JLTYP values for surface traction and edge load labels.**

Load Label	JLTYP	Load Label	JLTYP	Load Label	JLTYP
TRSHRNU	<b>510+j</b>	EDLDNU	<b>540+j</b>	EDTRANU	<b>570+j</b>
TRSHR1NU	<b>511</b>	EDLD1NU	<b>543</b>	EDTRANU	<b>573</b>
TRSHR2NU	<b>512</b>	EDLD2NU	<b>544</b>	EDTRANU	<b>574</b>
TRSHR3NU	<b>513</b>	EDLD3NU	<b>545</b>	EDTRANU	<b>575</b>
TRSHR4NU	<b>514</b>	EDLD4NU	<b>546</b>	EDTRANU	<b>576</b>
TRSHR5NU	<b>515</b>	EDNORNU	<b>550+j</b>	EDMOMNU	<b>580+j</b>
TRSHR6NU	<b>516</b>	EDNOR1NU	<b>553</b>	EDMOM1NU	<b>583</b>
TRVECNU	<b>520+j</b>	EDNOR2NU	<b>554</b>	EDMOM2NU	<b>584</b>
TRVEC1NU	<b>521</b>	EDNOR3NU	<b>555</b>	EDMOM3NU	<b>585</b>

<b>Load Label</b>	<b>JLTYP</b>	<b>Load Label</b>	<b>JLTYP</b>	<b>Load Label</b>	<b>JLTYP</b>
TRVEC2NU	<b>522</b>	EDNOR4NU	<b>556</b>	EDMOM4NU	<b>586</b>
TRVEC3NU	<b>523</b>	EDSHRNU	<b>560+j</b>		
TRVEC4NU	<b>524</b>	EDSHRNU	<b>563</b>		
TRVEC5NU	<b>525</b>	EDSHRNU	<b>564</b>		
TRVEC6NU	<b>526</b>	EDSHRNU	<b>565</b>		
		EDSHRNU	<b>566</b>		

**SNAME**

Surface name for a surface-based load definition. For an element-based or edge-based load the surface name is passed in as blank.

**1.1.49      UTRS: User subroutine to define a reduced time shift function for a viscoelastic material.**

**Product:** Abaqus/Standard

## References

---

- “Time domain viscoelasticity,” Section 19.7.1 of the Abaqus Analysis User’s Manual
- \*TRS
- \*VISCOELASTIC
- “Transient thermal loading of a viscoelastic slab,” Section 3.1.2 of the Abaqus Benchmarks Manual

## Overview

---

User subroutine **UTRS**:

- can be used to define a temperature-time shift for a time domain viscoelastic analysis;
- will be called for all material points of elements for which a user-defined shift function is specified to define the time-temperature correspondence as part of the viscoelastic material definition;
- will be called before user subroutine **UMATHT** and/or user subroutine **HETVAL** if either or both are to be used with **UTRS** in a fully coupled temperature-displacement analysis;
- can use and update solution-dependent state variables; and
- can be used in conjunction with user subroutine **USDFLD** to redefine any field variables before they are passed in.

## User subroutine interface

---

```

SUBROUTINE UTRS (SHIFT, TEMP, DTEMP, TIME, DTIME, PREDEF, DPRED,
1 STATEV, CMNAME, COORDS)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME
DIMENSION SHIFT(2), TIME(2), PREDEF(1), DPRED(1), STATEV(1),
1 COORDS(1)
C
user coding to define SHIFT(1) and SHIFT(2)
RETURN
END

```

**Variable to be defined**

---

**SHIFT**

An array of length two that defines the shift function,  $A$  ( $A > 0$ ), at this point. **SHIFT (1)** defines the shift function at the beginning of the increment, and **SHIFT (2)** defines the shift function at the end of the increment. Abaqus/Standard will apply an averaging scheme to these values that assumes that the natural logarithm of the shift function can be approximated by a linear function over the increment.

If either element of **SHIFT** is found to be less than or equal to zero, the analysis will terminate with an error message.

**Variable that can be updated**

---

**STATEV**

An array containing the solution-dependent state variables at this point. This array will be passed in containing the values of these variables at the start of the increment unless they are updated in user subroutines **USDFLD** or **UEXPAN**, in which case the updated values are passed in. If any of the solution-dependent state variables are being used in conjunction with the viscoelastic behavior, they must be updated in this subroutine to their values at the end of the increment.

**Variables passed in for information**

---

**TEMP**

Temperature at the end of the increment.

**DTEMP**

Increment of temperature during the time increment.

**PREFDEF**

An array containing the values of all of the user-specified field variables at this point at the end of the increment (initial values at the beginning of the analysis and current values during the analysis).

**DPRED**

An array containing the increments of all of the predefined field variables during the time increment.

**TIME (1)**

Value of step time at the end of the current increment.

**TIME (2)**

Value of total time at the end of the current increment.

**DTIME**

Time increment. If this subroutine is called during a procedure such as a static analysis in which the viscoelastic effects will not be taken into account, this variable is passed in as zero.

**CMNAME**

User-specified material name, left justified.

**COORDS**

An array containing the coordinates of the material point. These are the current coordinates if geometric nonlinearity is accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.



## 1.1.50 UVARM: User subroutine to generate element output.

**Product:** Abaqus/Standard

### References

---

- “Obtaining material point information in an Abaqus/Standard analysis,” Section 2.1.6
- \*USER OUTPUT VARIABLES
- “**UVARM**,” Section 4.1.26 of the Abaqus Verification Manual

### Overview

---

User subroutine **UVARM**:

- will be called at all material calculation points of elements for which the material definition includes the specification of user-defined output variables;
- may be called multiple times for each material point in an increment, as Abaqus/Standard iterates to a converged solution;
- will be called for each increment in a step;
- allows you to define output quantities that are functions of any of the available integration point quantities listed in the Output Variable Identifiers table (“Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual);
- allows you to define the material directions as output variables;
- can be used for gasket elements;
- can call utility routine **GETVRM** to access material point data;
- cannot be used with linear perturbation procedures; and
- cannot be updated in the zero increment.

### Accessing material point data

---

You are provided with access to the values of the material point quantities through the utility routine **GETVRM** described in “Obtaining material point information in an Abaqus/Standard analysis,” Section 2.1.6. In a nonlinear analysis values returned will correspond to the current solution iteration, representing a converged solution only at the final iteration for each increment. The values of the material point data are recovered in the arrays **ARRAY**, **JARRAY**, and **FLGRAY** for floating point, integer, and character data, respectively. Floating point data are recovered as double-precision data.

### Using user-defined output variables

---

The output identifier for the user-defined output quantities is UVARM. Individual components are accessed with **UVARM $n$** , where  $n = 1, 2, \dots, \text{NUVARM}$ . You must specify the number of user-defined output variables, **NUVARM**, for a given material to allocate space at each material calculation point for

each variable. The user-defined output variables are available for both printed and results file output and are written to the output database and restart files for contouring, printing, and  $X-Y$  plotting in Abaqus/CAE. Any number of user-defined output variables can be used.

**Input File Usage:** \*USER OUTPUT VARIABLES  
NUVARM

## Output precision

The data are provided in double precision for output to the data (.dat) and results (.fil) files and are written to the output database (.odb) file in single precision. Because the user provides **UVARM** output variables in double precision, numeric overflow errors related to output to the output database file may occur in cases where the output results exceed the capacity for single-precision representation even when no overflow errors occur in **UVARM**.

---

## User subroutine interface

```

SUBROUTINE UVARM(UVAR,DIRECT,T,TIME,DTIME,CNAME,ORNAME,
1 NUVARM,NOEL,NPT,LAYER,KSPT,KSTEP,KINC,NDI,NSHR,COORD,
2 JMAC,JMATYP,MATLAYO,LACCFLA)
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CNAME,ORNAME
CHARACTER*3 FLGRAY(15)
DIMENSION UVAR(NUVARM),DIRECT(3,3),T(3,3),TIME(2)
DIMENSION ARRAY(15),JARRAY(15),JMAC(*),JMATYP(*),COORD(*)

C      The dimensions of the variables FLGRAY, ARRAY and JARRAY
C      must be set equal to or greater than 15.

```

*user coding to define UVAR*

```

RETURN
END

```

---

## Variable to be defined

### UVAR (NUVARM)

An array containing the user-defined output variables. These are passed in as the values at the beginning of the increment and must be returned as the values at the end of the increment.

---

## Variables passed in for information

**DIRECT(3,3)**

An array containing the direction cosines of the material directions in terms of the global basis directions. **DIRECT(1,1)**, **DIRECT(2,1)**, **DIRECT(3,1)** give the (1, 2, 3) components of the first material direction; **DIRECT(1,2)**, **DIRECT(2,2)**, **DIRECT(3,2)** give the second material direction, etc. For shell and membrane elements the first two directions are in the plane of the element and the third direction is the normal. This information is not available for beam and truss elements.

**T(3,3)**

An array containing the direction cosines of the material orientation components relative to the element basis directions. This is the orientation that defines the material directions (**DIRECT**) in terms of the element basis directions. For continuum elements **T** and **DIRECT** are identical. For shell and membrane elements  $\mathbf{T}(1,1) = \cos \theta$ ,  $\mathbf{T}(1,2) = -\sin \theta$ ,  $\mathbf{T}(2,1) = \sin \theta$ ,  $\mathbf{T}(2,2) = \cos \theta$ ,  $\mathbf{T}(3,3) = 1.0$ , and all other components are zero, where  $\theta$  is the counterclockwise rotation around the normal vector that defines the orientation. If no orientation is used, **T** is an identity matrix. Orientation is not available for beam and truss elements.

**TIME(1)**

Value of step time at the end of the current increment.

**TIME(2)**

Value of total time at the end of the current increment.

**DTIME**

Time increment.

**CMNAME**

User-specified material name, left justified.

**ORNAME**

User-specified local orientation name, left justified.

**NUVARM**

User-specified number of user-defined output variables.

**NOEL**

Element number.

**NPT**

Integration point number.

**LAYER**

Layer number (for composite shells and layered solids).

**KSPT**

Section point number within the current layer.

**KSTEP**

Step number.

**KINC**

Increment number.

**NDI**

Number of direct stress components at this point.

**NSHR**

Number of shear stress components at this point.

**COORD**

Coordinates at this material point.

**JMAC**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**JMATYP**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**MATLayo**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**LACCFLA**

Variable that must be passed into the **GETVRM** utility routine to access an output variable.

**Example: Calculation of stress relative to shift tensor**

---

Below is an example of user subroutine **UVARM**. The subroutine calculates the position of the current state of stress relative to the center of the yield surface for the kinematic hardening plasticity model by subtracting the kinematic shift tensor,  $\alpha$ , from the stress tensor,  $\sigma$ . See “Metal plasticity models,” Section 4.3.1 of the Abaqus Theory Manual, for additional details.

```
SUBROUTINE UVARM(UVAR,DIRECT,T,TIME,DTIME,CMNAME,ORNAME,
1 NUVARM,NOEL,NPT,LAYER,KSPT,KSTEP,KINC,NDI,NSHR,COORD,
2 JMAC,JMATYP,MATLayo,LACCFLA)
C
INCLUDE 'ABA_PARAM.INC'
C
CHARACTER*80 CMNAME,ORNAME
CHARACTER*3 FLGRAY(15)
DIMENSION UVAR(NUVARM),DIRECT(3,3),T(3,3),TIME(2)
```

```

DIMENSION ARRAY(15),JARRAY(15),JMAC(*),JMATTYP(*),COORD(*)
C
C Error counter:
JERROR = 0
C Stress tensor:
CALL GETVRM('S',ARRAY,JARRAY,FLGRAY,JRCD,JMAC,JMATTYP,
1 MATLATO,LACCFLA)
JERROR = JERROR + JRCD
UVAR(1) = ARRAY(1)
UVAR(2) = ARRAY(2)
UVAR(3) = ARRAY(3)
UVAR(4) = ARRAY(4)
UVAR(5) = ARRAY(5)
UVAR(6) = ARRAY(6)
C Kinematic shift tensor:
CALL GETVRM('ALPHA',ARRAY,JARRAY,FLGRAY,JRCD,JMAC,JMATTYP,
1 MATLATO,LACCFLA)
JERROR = JERROR + JRCD
C Calculate the position relative to the center of the
C yield surface:
UVAR(1) = UVAR(1) - ARRAY(1)
UVAR(2) = UVAR(2) - ARRAY(2)
UVAR(3) = UVAR(3) - ARRAY(3)
UVAR(4) = UVAR(4) - ARRAY(4)
UVAR(5) = UVAR(5) - ARRAY(5)
UVAR(6) = UVAR(6) - ARRAY(6)
C If error, write comment to .DAT file:
IF(JERROR.NE.0)THEN
    WRITE(6,*) 'REQUEST ERROR IN UVARM FOR ELEMENT NUMBER ',
1      NOEL,'INTEGRATION POINT NUMBER ',NPT
ENDIF
RETURN
END

```



## 1.1.51 UWAVE: User subroutine to define wave kinematics for an Abaqus/Aqua analysis.

**Product:** Abaqus/Aqua

### References

---

- “Abaqus/Aqua analysis,” Section 6.11.1 of the Abaqus Analysis User’s Manual
- \*WAVE

### Overview

---

User subroutine **UWAVE**:

- will be called at each load integration point for which an Abaqus/Aqua load is specified and a user-defined gravity wave is specified;
- can be used to define the wave kinematics (fluid velocity and acceleration, dynamic pressure, vertical gradient of the dynamic pressure, and the instantaneous fluid surface elevation) as a function of time and space; and
- for stochastic analysis, can be used to determine when during the analysis the current configuration should be retained as the intermediate configuration upon which the wave kinematics are based.

### User subroutine interface

---

```

SUBROUTINE UWAVE (V,A,PDYN,DPDYNDZ,SURF,LPDYN
1 LRECOMPUTE,LUPLOCAL,LUPGLOBAL,
2 LSURF,NDIM,XCUR,XINTERMED,
3 GRAV,DENSITY,ELEV,B,ELEVS,
4 SEED,NSPECTRUM,FREQWAMP,
5 TIME,DTIME,NOEL,NPT,KSTEP,KINC)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION V(NDIM),A(NDIM),XCUR(NDIM),XINTERMED(NDIM),
1 FREQWAMP(2,NSPECTRUM),TIME(2)

```

*user coding to define V, A, PDYN, DPDYNDZ, SURF  
and, if necessary, LUPGLOBAL and LUPLOCAL*

```

RETURN
END

```

## Variables to be defined

---

### When LSURF=0

#### V (NDIM)

The total fluid velocity at the current load integration location. This array is passed into **UWAVE** as the steady current velocity. The array should be updated as the sum of the steady current velocity and the velocity contribution from the user-defined wave theory.

#### A (NDIM)

The fluid acceleration at the current load integration location.

#### PDYN

The dynamic pressure contribution to the total pressure. This variable is needed only for buoyancy loads. The total pressure at a location below the instantaneous surface elevation is the sum of the atmospheric pressure, the hydrostatic pressure measured to the mean fluid elevation, and the dynamic pressure. See “Airy wave theory,” Section 6.2.2 of the Abaqus Theory Manual, and “Stokes wave theory,” Section 6.2.3 of the Abaqus Theory Manual, for definitions of the dynamic pressure for Airy and Stokes waves, respectively.

#### DPDYNDZ

The gradient of the dynamic pressure in the vertical direction. This variable is needed only for buoyancy loads.

### When LSURF=1

#### SURF

The vertical coordinate of the instantaneous fluid surface corresponding to the horizontal position of the load integration point (given in **XCUR**). If the current location of the load integration point is above the instantaneous surface elevation, no fluid loads will be applied.

### Only in an analysis with stochastic wave kinematics based on an intermediate configuration

#### LUPLOCAL

Flag to determine if the intermediate configuration will be updated for this element. This flag can be set only when **LRECOMPUTE**=1. Return **LUPLOCAL** as 0 (default) to indicate that the intermediate configuration should not be updated. Return **LUPLOCAL** as 1 if the intermediate configuration should be updated for this element. The intermediate configuration is stored on an element-by-element basis. Therefore, all integration points for a given element will have their intermediate configuration updated if an update is requested at any one integration point on the element.

#### LUPGLOBAL

Flag to determine if the intermediate configuration will be updated for all elements. This flag can be set only when **LRECOMPUTE**=1. Return **LUPGLOBAL** as 0 (default) to indicate that the intermediate configuration should not be updated. Return **LUPGLOBAL** as 1 if the intermediate configuration should be updated for all elements with Abaqus/Aqua loads.

## Variables passed in for information

---

### **LRECOMPUTE**

For stochastic analysis **LRECOMPUTE**=1 indicates that an update to the intermediate configuration is permitted during this call to user subroutine **UWAVE**. The local and global update flags must be set accordingly. If the intermediate configuration is to be updated, the local update flag **LUPLOCAL** or the global update flag **LUPGLOBAL** must be set to 1. When **LRECOMPUTE**=1 and the intermediate configuration needs to be updated, the user subroutine should recompute all wave kinematics information based on the new intermediate configuration. For nonstochastic analysis this flag is always set to 0.

### **LPDYN**

**LPDYN**=1 indicates that only the dynamic pressure and its gradient need to be calculated (i.e., buoyancy loads). **LPDYN**=0 indicates that only the fluid velocity and acceleration need to be calculated (i.e., drag or inertia loads).

### **LSURF**

**LSURF**=1 indicates that subroutine **UWAVE** only needs to return the instantaneous fluid surface elevation. When **LSURF**=1, no velocity, acceleration, or dynamic pressure needs to be calculated. **LSURF**=0 indicates that the instantaneous fluid surface elevation **SURF** is not needed.

### **NDIM**

Two or three, indicating that the analysis is in two or three dimensions. The vertical direction is the global *y*-direction in two-dimensional analysis and the global *z*-direction in three-dimensional analysis.

### **XCUR (NDIM)**

An array containing the current coordinates of the load integration point.

### **XINTERMED (NDIM)**

An array containing the intermediate configuration coordinates of the load integration point. For nonstochastic analysis this array is not used. In a stochastic analysis the wave field is based upon this configuration. At the beginning of each load increment the **LRECOMPUTE** flag is set to 1 to prompt you for update action. If the intermediate configuration should be replaced by the current configuration, the flag **LUPLOCAL** should be set to 1 to update the intermediate configuration for this element only or the flag **LUPGLOBAL** should be set to 1 to update the intermediate configuration for all elements that have Abaqus/Aqua loading. At the beginning of the analysis the intermediate configuration is the reference configuration.

### **GRAV**

The user-specified gravitational constant in the fluid variable definition.

### **DENSITY**

The user-specified fluid mass density in the fluid variable definition.

**ELEV<sub>B</sub>**

The user-specified elevation of the seabed in the fluid variable definition.

**ELEV<sub>S</sub>**

The user-specified elevation of the still fluid level in the fluid variable definition.

**SEED**

For stochastic analysis the user-specified random number seed in the gravity wave definition.

**NSPECTRUM**

For stochastic analysis the number of user-specified frequency versus wave amplitude pairs in the gravity wave definition, used to define the wave spectrum.

**FREQWAMP (1 , NSPECTRUM)**

For stochastic analysis the frequency values used to define the wave spectrum.

**FREQWAMP (2 , NSPECTRUM)**

For stochastic analysis the wave amplitude values used to define the wave spectrum.

**TIME (1)**

Value of step time at the end of the current increment.

**TIME (2)**

Value of total time at the end of the current increment.

**DTIME**

Time increment.

**NOEL**

Element number.

**NPT**

Load integration point number. All line elements use full integration for the application of external loads. For distributed loads applied to the ends of the element, **NPT** corresponds to the end number of the element.

**KSTEP**

Step number.

**KINC**

Increment number.

## 1.1.52 VOIDRI: User subroutine to define initial void ratios.

**Product:** Abaqus/Standard

### References

---

- “Initial conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.2.1 of the Abaqus Analysis User’s Manual
- “Coupled pore fluid diffusion and stress analysis,” Section 6.8.1 of the Abaqus Analysis User’s Manual
- \*INITIAL CONDITIONS

### Overview

---

User subroutine **VOIDRI**:

- will be called to define initial void ratio values at material calculation points of continuum elements (see Part VI, “Elements,” of the Abaqus Analysis User’s Manual) in a porous medium whenever a user-defined initial condition on void ratio is specified; and
- can be used to define initial void ratio values as functions of material point coordinates and/or element numbers.

### User subroutine interface

---

```
SUBROUTINE VOIDRI (EZERO, COORDS, NOEL)
C
INCLUDE 'ABA_PARAM.INC'
C
DIMENSION COORDS (3)
C
```

*user coding to define EZERO*

```
RETURN
END
```

### Variable to be defined

---

#### EZERO

Initial void ratio.

---

## **Variables passed in for information**

### **COORDS**

An array containing the current coordinates of this point.

### **NOEL**

Element number.

## 1.2 Abaqus/Explicit subroutines

- “VDISP,” Section 1.2.1
- “VDLOAD,” Section 1.2.2
- “VFABRIC,” Section 1.2.3
- “VFRIC,” Section 1.2.4
- “VFRIC\_COEF,” Section 1.2.5
- “VFRICITION,” Section 1.2.6
- “VUAMP,” Section 1.2.7
- “VUANISOHYPER\_INV,” Section 1.2.8
- “VUANISOHYPER\_STRAIN,” Section 1.2.9
- “VUEL,” Section 1.2.10
- “VUFIELD,” Section 1.2.11
- “VUFLUIDEXCH,” Section 1.2.12
- “VUFLUIDEXCHEFFAREA,” Section 1.2.13
- “VUHARD,” Section 1.2.14
- “VUINTER,” Section 1.2.15
- “VUINTERACTION,” Section 1.2.16
- “VUMAT,” Section 1.2.17
- “VUSDFLD,” Section 1.2.18
- “VUTRS,” Section 1.2.19
- “VUVISCOSITY,” Section 1.2.20



## 1.2.1 VDISP: User subroutine to specify prescribed boundary conditions.

**Product:** Abaqus/Explicit

### References

---

- “Boundary conditions in Abaqus/Standard and Abaqus/Explicit,” Section 30.3.1 of the Abaqus Analysis User’s Manual
- \*BOUNDARY
- “**VDISP**,” Section 4.1.28 of the Abaqus Verification Manual

### Overview

---

User subroutine **VDISP**:

- can be used to prescribe translational and rotational boundary conditions;
- is called for all degrees of freedom listed in a user subroutine-defined boundary condition;
- defines the magnitudes of the specified type of the associated boundary condition; and
- can be called for blocks of nodes for which the boundary conditions are defined in the subroutine.

### Initialization

---

At the beginning of a step user subroutine **VDISP** is called twice to establish all required initial conditions.

The first call to user subroutine **VDISP** is made to establish the startup mean velocity, which is indicated by the passing of a step time value of  $-dt$  into the subroutine, where  $dt$  is the current time increment. If displacement is prescribed, the returned variable, **rval**, corresponds to the displacement at  $-dt$ . In this case **rval** should be set equal to  $u_o - v_o dt + \frac{1}{2}a_o dt^2$ , where  $u_o$ ,  $v_o$  and  $a_o$  are the initial displacement, velocity, and acceleration, respectively. If velocity is prescribed, the returned variable corresponds to the mean velocity at  $-dt/2$ . In this case **rval** should be set equal to  $v_o - \frac{1}{2}a_o dt$ . If acceleration is prescribed, the returned variable corresponds to the acceleration at  $-dt$ . In this case **rval** should be set equal to  $\frac{1}{dt}v_o - \frac{1}{2}a_o$ . The initial displacement and velocity are passed into the user subroutine in the arrays **u** and **v**, respectively.

The second call to user subroutine **VDISP** is made to establish the initial acceleration, which is indicated by the passing of a step time value of zero into the subroutine. If displacement is prescribed, the returned variable should be set equal to the displacement at  $dt$ . If velocity is prescribed, the returned variable should be set equal to the mean velocity at  $dt/2$ . If acceleration is prescribed, the returned variable should be set equal to the acceleration at zero step time.

### Time incrementation

---

During time incrementation user subroutine **VDISP** is called once for each increment to establish all required prescribed conditions.

If displacement is prescribed, the returned variable should be set equal to the displacement at **stepTime+dtNext**, where **stepTime** is the step time and **dtNext** is the next time increment. If velocity is prescribed, the returned variable should be set equal to the mean velocity at **stepTime+dtNext/2**. If acceleration is prescribed, the returned variable should be set equal to the acceleration at **stepTime**.

## User subroutine interface

---

```

      subroutine vdisp(
c Read only variables -
  1   nblock, nDof, nCoord, kstep, kinc,
  2   stepTime, totalTime, dtNext, dt,
  3   cbname, jBCType, jDof, jNodeUid, amp,
  4   coordNp, u, v, a, rf, rmass, rotaryI,
c Write only variable -
  5   rval )
c
      include 'vaba_param.inc'
c
      character*80 cbname
      dimension jDof(nDof), jNodeUid(nblock),
  1          amp(nblock), coordNp(nCoord,nblock),
  2          u(nDof,nblock), v(nDof,nblock), a(nDof,nblock),
  3          rf(nDof,nblock), rmass(nblock), rotaryI(3,3,nblock),
  4          rval(nDof,nblock)
c
      do 100 k = 1, nblock
      do 100 j = 1, nDof
          if( jDof(j) .gt. 0 ) then
              user coding to define rval(j, k)
          end if
  100 continue
c
      return
end

```

## Variable to be defined

---

**rval(nDof, nblock)**

Values of the prescribed variable for degrees of freedom 1–6 (translation and rotation) at the nodes. The variable can be displacement, velocity, or acceleration, depending on the type specified in the associated boundary condition. The variable type is indicated by **jBCType**.

---

**Variables passed in for information****nblock**

Number of nodal points to be processed in this call to **VDISP**.

**nDof**

Number of degrees of freedom (equals 6).

**nCoord**

Number of coordinate components (equals 3).

**kstep**

Step number.

**kinc**

Increment number.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**dtNext**

Next time increment size.

**dt**

Current time increment size.

**cbname**

User-specified name corresponding to the associated boundary condition.

**jBCType**

Indicator for type of prescribed variable: 0 for displacement, 1 for velocity, and 2 for acceleration.

**jDof (nDof)**

Indicator for prescribed degrees of freedom. The values given by **rval(j,k)** are prescribed only if **jDof(j)** equals 1.

**jNodeUid (nblock)**

Node numbers.

**amp (nblock)**

Amplitude values corresponding to the associated amplitude functions. These values are passed in for information only and will not contribute to the values of the prescribed variable automatically.

**coordNp (nCoord, nblock)**

Nodal point coordinates.

**u (nDof, nblock)**

Nodal point displacements at **stepTime**. All translations are included if one or more translational degrees of freedom are prescribed. All rotations are included if one or more rotational degrees of freedom are prescribed.

**v (nDof, nblock)**

Nodal point velocities at zero step time during initialization or nodal point mean velocities at **stepTime-dt/2** during time incrementation. All translational velocities are included if one or more translational degrees of freedom are prescribed. All angular velocities are included if one or more rotational degrees of freedom are prescribed.

**a (nDof, nblock)**

Nodal point accelerations at **stepTime** before the boundary condition is prescribed. All translational accelerations are included if one or more translational degrees of freedom are prescribed. All angular accelerations are included if one or more rotational degrees of freedom are prescribed.

**rf (nDof, nblock)**

Nodal point reaction at **stepTime-dt**. All reaction forces are included if one or more translational degrees of freedom are prescribed. All reaction moments are included if one or more rotational degrees of freedom are prescribed.

**rmass (nblock)**

Nodal point masses.

**rotaryI (3, 3, nblock)**

Nodal point rotary inertia.

---

**Example: Imposition of acceleration on a rigid body with nonzero initial velocity**


---

In this example a sinusoidal acceleration is imposed on the reference node of a rigid body. Nonzero initial velocity is also specified for the rigid body. User subroutine **VDISP** given below illustrates how the return value array is to be computed for different phases of the solution. The analysis results show that both the initial velocity and acceleration are correctly specified.

**Input file**

```
*HEADING
  Test VDISP with S4R element
*NODE, NSET=NALL
  1,
  2, 2., 0.
  3, 0., 2.
  4, 2., 2.
```

```

 9, 1., 1., 0.
*ELEMENT, TYPE=S4R, ELSET=SHELL
 10, 1,2,4,3
*SHELL SECTION, ELSET=SHELL, MATERIAL=ELSHELL
 2.000000e-02,      3
*MATERIAL, NAME=ELSHELL
*DENSITY
7850.0,
*ELASTIC
 2.500000e+11,   3.000000e-01
*RIGID BODY, REF NODE=9, ELSET=SHELL
*INITIAL CONDITIONS, Type=VELOCITY
 9, 1, 0.4
*STEP
*DYNAMIC, EXPLICIT, DIRECT USER CONTROL
 0.01, 0.8
*BOUNDARY, USER, TYPE=ACCELERATION
 9, 1
*OUTPUT,HISTORY, TIME INTERVAL=0.01, OP=NEW
*NODE OUTPUT, NSET=NALL
 U, V, A
*END STEP

```

#### User subroutine

```

    subroutine vdisp(
c Read only variables -
    * nblock, nDof, nCoord, kstep, kinc,
    * stepTime, totalTime, dtNext, dt,
    * cbname, jBCType, jDof, jNodeUid, amp,
    * coordNp, u, v, a, rf, rmass, rotaryI,
c Write only variable -
    * rval )
c
    include 'vaba_param.inc'
    parameter( zero = 0.d0, half = 0.5d0, one = 1.d0 )
c
    character*80 cbname
    dimension jDof(nDof), jNodeUid(nblock),
    * amp(nblock), coordNp(nCoord,nblock),
    * u(nDof,nblock), v(nDof,nblock), a(nDof,nblock),
    * rf(nDof,nblock), rmass(nblock),
    * rotaryI(3,3,nblock), rval(nDof,nblock)

```

```
c
c      Impose acceleration
c
c      if( jBCType .eq. 2 ) then
c
c          if( stepTime .lt. zero ) then
c
c              Initialization 1
c
c              a0 = zero
c              do 310 k=1, nblock
c                  do 310 j=1, nDof
c                      if ( jDof(j) .gt. 0 ) then
c                          v0 = v(j,k)
c                          rval(j,k) = v0/dt - a0*half
c                      end if
c 310          continue
c
c          else if( stepTime .eq. zero ) then
c
c              Initialization 2
c
c              a0 = zero
c              do 320 k=1, nblock
c                  do 320 j=1, nDof
c                      if ( jDof(j) .gt. 0 ) then
c                          rval(j,k) = a0
c                      end if
c 320          continue
c
c          else
c
c              Time incrementation
c
c              amplitude = 2.0
c              period = 0.8
c              twopi = 6.2831853d0
c
c              do 350 k=1, nblock
c                  do 350 j=1, nDof
c                      if ( jDof(j) .gt. 0 ) then
c                          rval(j,k) = amplitude*
```

```
*                      sin( twopi*stepTime / period )
      end if
350      continue
      end if
end if
c
return
end
```



## 1.2.2 VDLOAD: User subroutine to specify nonuniform distributed loads.

**Product:** Abaqus/Explicit

### References

---

- “Applying loads: overview,” Section 30.4.1 of the Abaqus Analysis User’s Manual
- “Distributed loads,” Section 30.4.3 of the Abaqus Analysis User’s Manual
- \*DLOAD
- \*DSLOAD
- “Deformation of a sandwich plate under CONWEP blast loading,” Section 9.1.8 of the Abaqus Example Problems Manual

### Overview

---

User subroutine **VDLOAD**:

- can be used to define the variation of the distributed load magnitude as a function of position, time, velocity, etc. for a group of points, each of which appears in an element-based or surface-based nonuniform load definition;
- will be called for load integration points associated with each nonuniform load definition including PENU and PINU loads applicable for pipe elements;
- does not make available the current value of the nonuniform distributed loads for file output purposes; and
- recognizes an amplitude reference (“Amplitude curves,” Section 30.1.2 of the Abaqus Analysis User’s Manual) if it appears with the associated nonuniform load definition.

### User subroutine interface

---

```

        subroutine vdload (
C Read only (unmodifiable) variables -
        1 nblock, ndim, stepTime, totalTime,
        2 amplitude, curCoords, velocity, dirCos, jltyp, sname,
C Write only (modifiable) variable -
        1 value )
C
        include 'vaba_param.inc'
C
        dimension curCoords(nblock,ndim), velocity(nblock,ndim),
        1 dirCos(nblock,ndim,ndim), value(nblock)
        character*80 sname
C

```

```

do 100 km = 1, nblock
  user coding to define value

100 continue

return
end

```

**Variable to be defined**

---

**value (nblock)**

Magnitude of the distributed load. Units are  $\text{FL}^{-2}$  for surface loads,  $\text{FL}^{-3}$  for body forces.

**Variables passed in for information**

---

**nblock**

Number of points to be processed in this call to **VDLOAD**.

**ndim**

Number of coordinate directions: 2 for two-dimensional models, 3 for three-dimensional models. The model will be considered three-dimensional if any three-dimensional elements are defined (including SPRINGA elements).

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime - stepTime**.

**amplitude**

Current value of the amplitude referenced for this load (set to unity if no amplitude is referenced). You must multiply the load by the current amplitude value within the user subroutine if the amplitude is required.

**curCoords (nblock, ndim)**

Current coordinates of each point for which the load is to be calculated.

**velocity (nblock, ndim)**

Current velocity of each point for which the load is to be calculated.

**dirCos (nblock, ndim, ndim)**

Current orientation of the face, edge, pipe, or beam for pressure type loads (not applicable for body force type loads). The second dimension indicates the vector, and the third dimension indicates the components of that vector. For faces (pressures on three-dimensional continuum, shell, and membrane elements), the first and second vectors are the local directions in the plane of the surface and the third

vector is the normal to the face, as defined in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual. For solid elements the normal points inward, which is the opposite of what is defined in the conventions; for shell elements the normal definition is consistent with the defined conventions. For edges (pressures on two-dimensional continuum elements and two-dimensional beams and pipes), the first vector is the normal to the edge, the second vector is the tangent to the edge, and, if **ndim**=3, the third vector will be a unit normal in the out-of-plane direction. For three-dimensional beam and pipe elements, the first and second vectors are the local axes ( $\mathbf{n}_1, \mathbf{n}_2$ ) and the third vector is the tangent vector ( $\mathbf{t}$ ), as defined in “Beam element cross-section orientation,” Section 26.3.4 of the Abaqus Analysis User’s Manual.

### **j1typ**

Key that identifies the distributed load type. The load type may be a body force, a surface-based load, or an element-based surface load. For element-based surface loads, this variable identifies the element face for which this call to **VDLOAD** is being made. See Part VI, “Elements,” of the Abaqus Analysis User’s Manual for element load type identification. This information is useful when several different nonuniform distributed loads are being imposed on an element at the same time. The key is as follows:

<b>J1type</b>	<b>Load type</b>
0	Surface-based load
1	BXNU
2	BYNU
3	BZNU
20	PNU
21	P1NU
22	P2NU
23	P3NU
24	P4NU
25	P5NU
26	P6NU
27	PINU
28	PENU
41	PXNU
42	PYNU
43	PZNU

## VDLOAD

### **sname**

Surface name for a surface-based load definition (**JLTYP**=0). For a body force or an element-based load the surface name is passed in as a blank.

### 1.2.3 VFABRIC: User subroutine to define fabric material behavior.

**Product:** Abaqus/Explicit

*WARNING: The use of this user subroutine generally requires considerable expertise. You are cautioned that the implementation of any realistic constitutive model requires extensive development and testing. Initial testing on a single-element model with prescribed traction loading is strongly recommended.*

#### References

---

- “Fabric material behavior,” Section 20.4.1 of the Abaqus Analysis User’s Manual
- \*FABRIC

#### Overview

---

User subroutine **VFABRIC**:

- is used to define the mechanical constitutive behavior of a fabric material in the plane of the fabric;
- is valid for materials that exhibit two “structural” directions that may not be orthogonal to each other with deformation;
- is used to update the nominal fabric stresses for a given nominal fabric strain where the direct strains are defined as the nominal strain measured along the two yarn directions of the fabric and the engineering shear strain is defined as the drop in the angle between the two yarn directions going from the reference configuration to the current configuration;
- can be used with elements under plane stress conditions;
- will be called for blocks of material calculation points for which the material is defined in a user subroutine (“Material data definition,” Section 18.1.2 of the Abaqus Analysis User’s Manual);
- can use and update solution-dependent state variables;
- can use any field variables that are passed in; and
- cannot be used in an adiabatic analysis.

#### Component ordering in tensors

---

The component ordering depends upon whether the tensor is a “strain” variable or a “stress” variable.

#### Symmetric tensors

Tensors such as the strain and strain increment have four components, and tensors such as stress have three components, with the difference between the two sets of variables arising from the assumed plane stress condition. The component order with the arrays for these variables is listed in the table below:

Component	Strain	Stress
1	$\varepsilon_{11}$	$\sigma_{11}$
2	$\varepsilon_{22}$	$\sigma_{22}$
3	$\varepsilon_{33}$	$\sigma_{12}$
4	$\varepsilon_{12}$	

The shear strain components in user subroutine **VFABRIC** are stored as tensor components and not as engineering components.

### Initial calculations and checks

---

In the **datacheck** phase of the analysis Abaqus/Explicit calls user subroutine **VFABRIC** with a set of fictitious strains and a **totalTime** and **stepTime** that are both equal to 0.0. This step serves as a check on your constitutive relation and calculates the equivalent initial material properties, upon which the initial elastic wave speeds are computed.

### Orientation of the fabric yarn

---

In general, the yarn directions may not be orthogonal to each other in the reference configuration. You can specify these local directions with respect to the in-plane axes of an orthogonal orientation system at a material point. Both the local directions and the orthogonal system are defined together as a single orientation definition. If the local directions are not specified, these directions are assumed to match the in-plane axes of the orthogonal system. The local direction may not remain orthogonal with deformation. Abaqus updates the local directions with deformation and computes the nominal strains along these directions and the drop in angle between them (the fabric engineering shear strain). The constitutive behavior for the fabric defines the fabric nominal stresses as a function of the fabric strains. Abaqus converts these fabric stresses into the Cauchy stress and the resulting internal forces.

### Material point deletion

---

Material points that satisfy a user-defined failure criterion can be deleted from the model (see “User-defined mechanical material behavior,” Section 23.8.1 of the Abaqus Analysis User’s Manual). You must specify the state variable number controlling the element deletion flag when you allocate space for the solution-dependent state variables, as explained in “Fabric material behavior,” Section 20.4.1 of the Abaqus Analysis User’s Manual. The deletion state variable should be set to a value of one or zero in user subroutine **VFABRIC**. A value of one indicates that the material point is active, while a value of zero indicates that Abaqus/Explicit should delete the material point from the model by setting the stresses to zero. The structure of the block of material points passed to user subroutine **VFABRIC** remains unchanged during the analysis; deleted material points are not removed from the block. Abaqus/Explicit will pass zero stresses and strain increments for all deleted material points. Once a material point has been flagged as deleted, it cannot be reactivated.

---

User subroutine interface

```

        subroutine vfabric(
C Read only (unmodifiable) variables -
  1      nblock, ndim, npt, layer, kspt, kstep, kinc,
  2      nstatev, nfieldv, nprops,
  3      lOp, jElem, stepTime, totalTime, dt, cmname, coordMp,
  4      charLength, props, density, braidAngle, fabricStrain,
  5      fabricStrainInc,
  6      tempOld, fieldOld, fabricStressOld, stateOld,
  7      tempNew, fieldNew, enerIntern,
C Write only (modifiable) variables -
  8      fabricStressNew, stateNew, enerInelas )
C
C NOTE: In addition to the above "Write only" variables,
C the thickness direction component of fabricStrainInc
C i.e, fabricStrainInc(*,ndirStrain) may also be set by
C the user for changing thickness as a function
C of material in-plane state.
C
        include 'vaba_param.inc'
C
        parameter ( ndirStrain = 3, nshr = 1, ndirStress = 2)
C
C NOTE: The constants defined above are used for array
C dimensions below.
C
        dimension
*      jElem(nblock),
*      coordMp(nblock,ndim),
*      charLength(nblock),
*      props(nprops),
*      density(nblock),
*      braidAngle(nblock),
*      fabricStrain(nblock,ndirStrain+nshr),
*      strainFabricInc(nblock,ndirStrain+nshr),
*      tempOld(nblock),
*      fieldOld(nblock,nfieldv),
*      fabricStressOld(nblock,ndirStress+nshr),
*      stateOld(nblock,nstatev),
*      tempNew(nblock),

```

```

*      fieldNew(nblock,nfieldv),
*      fabricStressNew(nblock,ndirStress+nshr),
*      stateNew(nblock,nstatev),
*      enerIntern(nblock),
*      enerInelas(nblock)
*
*      character*80 cmname
C
do 100 km = 1,nblock
  user coding
100 continue

return
end

```

**Variables to be defined**

---

**fabricStressNew(nblock,ndirStress+nshr)**

Nominal fabric stress at each material point at the end of the increment. This nominal fabric stress can be requested as output variable SFABRIC.

**stateNew(nblock,nstatev)**

State variables at each material point at the end of the increment. You define the size of this array by allocating space for it (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information). This variable can be requested as output variable SDV.

**enerInelas(nblock)**

Total inelastic energy density at material points at the end of the increment. This variable can be requested as output variable ENER.

**Variable that can be updated**

---

**fabricStrainInc(\*,ndirStrain)**

Thickness direction strain increment. The thickness can be requested as output variable STH.

**Variables passed in for information**

---

**nblock**

Number of material points to be processed in this call to **VFABRIC**.

**ndim**

Two for a two-dimensional model and three for a three-dimensional model.

**npt**

Current integration point number.

**layer**

Current layer number in the case of a composite section.

**kspt**

Current material point number within the section.

**kStep**

Current Abaqus step number.

**kInc**

Increment number of the current Abaqus step.

**nstatev**

Number of user-defined state variables that are associated with this material type (you define this as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**lOp**

Integer flag indicating the computation that is expected. **lOp** = -2 indicates that the routine is being called to initialize the stresses corresponding to the initial strains, which can be large. **lOp** = -1 indicates that the routine is being called to update the stresses based on the instantaneous elastic response for a small “artificial” strain increment given. **lOp** = 0 indicates that this is an annealing process and you should reinitialize the internal state variables, **stateNew**, if necessary. The stresses will be set to zero by Abaqus. **lOp** = 1 indicates that the routine is being called to update the stresses and the state for a given strain increment.

**jElem(nblock)**

Array of element numbers.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime** - **stepTime**.

**dt**

Time increment size.

**cmname**

User-specified material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the character string “ABQ\_”. To avoid conflict, you should not use “ABQ\_” as the leading string for **cmname**.

**coordMp (nblock, \*)**

Material point coordinates. It is the midplane material point for shell elements and the centroid for beam elements.

**charLength (nblock)**

Characteristic element length. This is a typical length of a line across an element for a first-order element; it is half of the same typical length for a second-order element. For membranes and shells it is a characteristic length in the reference surface.

**props (nprops)**

User-supplied material properties.

**density (nblock)**

Current density at the material points in the midstep configuration. This value may be inaccurate in problems where the volumetric strain increment is very small. If an accurate value of the density is required in such cases, the analysis should be run in double precision. This value of the density is not affected by mass scaling.

**braidAngle (nblock)**

Angle in radians between the two yarn directions at the end of the increment.

**fabricStrain (nblock,ndirStrain+nshr)**

Total nominal strain in the fabric at the end of increment. This variable can be requested as output variable EFABRIC.

**fabricStrainInc (nblock,ndirStrain+nshr)**

Incremental nominal strain in the fabric.

**tempOld (nblock)**

Temperatures at each material point at the beginning of the increment.

**fieldOld (nblock,nfieldv)**

Values of the user-defined field variables at each material point at the beginning of the increment.

**fabricStressOld (nblock,ndirStress+nshr)**

Nominal fabric stress at each material point at the beginning of the increment.

**stateOld (nblock,nstatev)**

State variables at each material point at the beginning of the increment.

**tempNew(nblock)**

Temperatures at each material point at the end of the increment.

**fieldNew(nblock, nfieldv)**

Values of the user-defined field variables at each material point at the end of the increment.

**enerIntern(nblock)**

Internal energy per unit mass at each material point at the beginning of the increment.

**Example: Using more than one user-defined material model**

---

To use more than one user-defined fabric material model, the variable **cmname** can be tested for different fabric material names inside user subroutine **VFABRIC**, as illustrated below:

```
if (cmname(1:4) .eq. 'MAT1') then
    call VFABRIC_MAT1(argument_list)
else if (cmname(1:4) .eq. 'MAT2') then
    call VFABRIC_MAT2(argument_list)
end if
```

**VFABRIC\_MAT1** and **VFABRIC\_MAT2** are the actual fabric material user subroutines containing the constitutive material models for each material **MAT1** and **MAT2**, respectively. User subroutine **VFABRIC** merely acts as a directory here. The argument list can be the same as that used in subroutine **VFABRIC**. The material names must be in uppercase characters since **cmname** is passed in as an uppercase character string.

**Example: Influence of nonorthogonal material directions in highly anisotropic elastic material**

---

As an example of the coding of user subroutine **VFABRIC**, consider a simple elastic lamina material with highly anisotropic properties. For a fabric the material definitions need not remain orthogonal with deformation, whereas the directions do remain orthogonal for a built-in elastic material. The simple **VFABRIC** routine given below defines an elastic fabric and can be used to compare the fabric and the built-in elastic materials under different loading conditions.

The user subroutine would be coded as follows:

```
subroutine vfabric(
C Read only (unmodifiable)variables -
  1      nblock, ndim, npt, layer, kspt, kstep, kinc,
  2      nstatev, nfieldv, nprops,
  3      lOp, jElem, stepTime, totalTime, dt, cmname, coordMp,
  4      charLength, props, density, braidAngle, fabricStrain,
  5      fabricStrainInc,
  6      tempOld, fieldOld, fabricStressOld, stateOld,
  7      tempNew, fieldNew, enerIntern,
C Write only (modifiable) variables -
```

```
     8      fabricStressNew, stateNew, enerInelas )  
C  
C NOTE: In addition to the above "Write only" variables,  
C the thickness direction component of fabricStrainInc  
C i.e., fabricStrainInc(*,ndirStrain) may also be set by the user  
C for changing thickness as a function of material in-plane  
C state.  
C  
     include 'vaba_param.inc'  
C  
     parameter( ndirStrain = 3, nshr = 1, ndirStress = 2,  
              one = 1.d0, two = 2.d0 )  
C  
C NOTE: The constants defined above are used for array  
C dimensions and computation below.  
C  
     dimension  
     *      jElem(nblock),  
     *      coordMp(nblock,ndim),  
     *      charLength(nblock),  
     *      props(nprops),  
     *      density(nblock),  
     *      braidAngle(nblock),  
     *      fabricStrain(nblock,ndirStrain+nshr),  
     *      fabricStrainInc(nblock,ndirStrain+nshr),  
     *      tempOld(nblock),  
     *      fieldOld(nblock,nfieldv),  
     *      fabricStressOld(nblock,ndirStress+nshr),  
     *      stateOld(nblock,nstatev),  
     *      tempNew(nblock),  
     *      fieldNew(nblock,nfieldv),  
     *      fabricStressNew(nblock,ndirStress+nshr),  
     *      stateNew(nblock,nstatev),  
     *      enerIntern(nblock),  
     *      enerInelas(nblock)  
C  
     character*80 cmname  
C  
C  
C Read properties
```

```

E1      = props(1)
E2      = props(2)
xnu12  = props(3)
twiceG12 = two * props(4)
C
xnu21 = E2 * xnu12 / E1
C
C Let us assume:
xnu13 = xnu12
xnu23 = xnu21
C
xnu13OverE1 = xnu13/E1
xnu23OverE2 = xnu23/E2
C
fr = one / ( one - xnu12 * xnu21 )
D11 = E1 * fr
D22 = E2 * fr
D12 = E2 * xnu12 * fr
C
do k = 1 , nblock
C
C Update the stress
stressInc11 = D11 * fabricStrainInc(k,1)
*
*          + D12 * fabricStrainInc(k,2)
stressInc22 = D22 * fabricStrainInc(k,2)
*
*          + D12 * fabricStrainInc(k,1)
stressInc12 = twiceG12 *
*
*          fabricStrainInc(k,ndirStrain + 1)
C
fabricStressNew(k,1) = fabricStressOld(k,1)
*
*          + stressInc11
fabricStressNew(k,2) = fabricStressOld(k,2)
*
*          + stressInc22
C
C          shear stress
fabricStressNew(k,ndirStress+1) =
*
*          fabricStressOld(k,ndirStress+1) + stressInc12
C
C          Thickness direction strain
C          fabricStrainInc(k,ndirStrain) =
C          *          - ( xnu13OverE1 * stressInc11
C          *          + xnu23OverE2 * stressInc22

```

VFABRIC

```
C      *      )
C
end do

return
end
```

## 1.2.4 VFRIC: User subroutine to define frictional behavior for contact surfaces.

**Product:** Abaqus/Explicit

### References

---

- “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual
- \*FRICTION
- “**VFRIC**, **VFRIC\_COEF**, and **VFRICTION**,” Section 4.1.30 of the Abaqus Verification Manual

### Overview

---

User subroutine **VFRIC**:

- can be used to define the frictional behavior between contact pair surfaces;
- can be used when the classical Coulomb friction model is too restrictive and a more complex definition of shear transmission between contacting surfaces is required;
- must provide the entire definition of shear interaction between the contacting surfaces;
- can use and update solution-dependent state variables;
- cannot be used in conjunction with softened tangential surface behavior; and
- cannot be used with the general contact algorithm.

### Terminology

---

The use of user subroutine **VFRIC** requires familiarity with the following terminology.

#### Surface node numbers

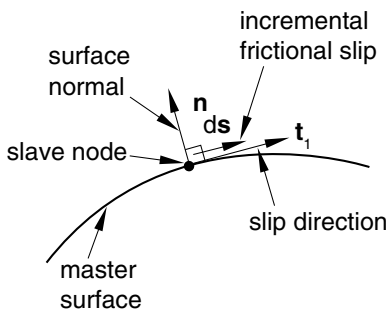
The “surface node number” refers to the position of a particular node in the list of nodes on the surface. For example, there are **nSlvNod** nodes on the slave surface. Number  $n, n = 1, 2, \dots, \text{nSlvNod}$ , is the surface node number of the  $n$ th node in this list; **jSlvUid**( $n$ ) is the user-defined global number of this node. An Abaqus/Explicit model can be defined in terms of an assembly of part instances (see “Defining an assembly,” Section 2.9.1 of the Abaqus Analysis User’s Manual). In such models a node number in **jSlvUid** is an internally generated node number. If the original node number and part instance name are required, call the utility routine **VGETPARTINFO** (see “Obtaining part information,” Section 2.1.5).

#### Contact points

The nodes on the slave surface that are in contact in the current time increment are defined as “contact points.” The number of contact points is passed into this subroutine as **nContact**. The array **jConSlvid(nContact)** gives the surface node numbers for the contact points.

## Local coordinate system

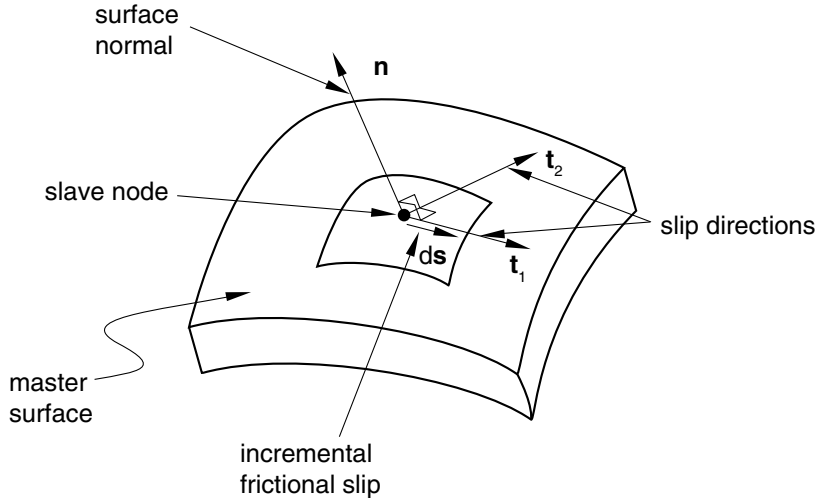
A local coordinate system is defined for each contact point to facilitate specification of frictional forces and incremental slips. The local 1-direction for both two-dimensional and three-dimensional contact is tangential to the master surface, and it is defined by  $t_1 = ds/|ds|$ , where  $ds$  is the incremental slip vector. The incremental slip vector used to define  $t_1$  corresponds to the incremental slip in the current time increment for penalty contact and the predicted incremental slip for kinematic contact. The master surface normal direction,  $n$ , is the local 2-direction for two-dimensional contact and the local 3-direction for three-dimensional contact. The local 2-direction for three-dimensional contact is given by  $t_2 = n \times t_1$ , which is also tangent to the master surface. The vectors are shown in Figure 1.2.4–1 and Figure 1.2.4–2. The direction cosines for  $t_1$  and  $n$  with respect to the global coordinate system are available in **dirCosT1** and **dirCosN**, respectively. In the case of zero incremental slip ( $|ds| = 0$ ) we choose an arbitrary direction for  $t_1$  that is orthogonal to the normal direction,  $n$ .



**Figure 1.2.4–1** Local coordinate system for two-dimensional contact with **VFRIC**.

## Frictional forces

You specify the frictional force, **fTangential**, at each contact point in local coordinates in this subroutine. The array **fTangential** is dimensioned such that only the tangential components can be specified. Any components of the frictional force that are not specified will remain equal to zero. For three-dimensional contact with isotropic friction, only the first component of the frictional force need be specified since the second component should be zero. A “stick force” at each contact point is provided in the array **fStickForce** to assist you in setting appropriate frictional force values. The stick force is the force required to prevent additional “plastic” slipping. The stick force at each contact point is provided as a scalar value as it would act in the direction opposite to  $t_1$ . The stick force is computed prior to calling user subroutine **VFRIC** by either the kinematic or the penalty contact algorithm. See “Contact constraint enforcement methods in Abaqus/Explicit,” Section 34.2.3 of the Abaqus Analysis User’s Manual, for descriptions of the kinematic and penalty contact algorithms and the user interface for choosing between them. The first component of the frictional force should be in the range between



**Figure 1.2.4–2** Local coordinate system for three-dimensional contact with **VFRIC**.

zero and minus the stick force value. Typically, the stick force will be positive and the first component of the applied frictional force will be negative, opposing the incremental slip. Penalty contact includes an elastic slip regime due to finite penalty stiffness, so occasionally, during recovery of elastic slip, the stick force will be negative, indicating that it is appropriate for the first component of the frictional force to be positive (i.e., acting in the same direction as the incremental slip). A noisy or unstable solution is likely to result if the first component of **fTangential** is set outside of the range between zero and negative the value of the stick force.

After user subroutine **VFRIC** is called, frictional forces that oppose the forces specified at the contact points are distributed to the master nodes. For balanced master-slave contact we then compute weighted averages of the frictional forces for both master-slave orientations. These forces are directly applied if the penalty contact algorithm is being used. If the kinematic contact algorithm is being used, the frictional forces are converted to acceleration corrections by dividing by the nodal masses.

### User subroutine interface

---

```

subroutine vfric(
C Write only -
  1 fTangential,
C Read/Write -
  2 statev,
C Read only -
  3 kStep, kInc, nContact, nFacNod, nSlvNod, nMstNod,
  4 nFricDir, nDir, nStateVar, nProps, nTemp, nPred, numDefTfv,
  5 jSlvUid,jMstUid, jConSlvid, jConMstid, timStep, timGlb,
```

```

6 dTimCur, surfInt, surfSlv, surfMst, lContType,
7 dSlipFric, fStickForce, fTangPrev, fNormal, frictionWork,
8 shape, coordSlv, coordMst, dirCosSl, dircosN, props,
9 areaSlv, tempSlv, preDefSlv, tempMst, preDefMst)

C
  include `vaba_param.inc'
C
  character*80 surfInt, surfSlv, surfMst
C
  dimension props(nProps), statev(nStateVar,nSlvNod),
1 dSlipFric(nDir,nContact),
2 fTangential(nFricDir,nContact),
3 fTangPrev(nDir,nContact),
4 fStickForce(nContact), areaSlv(nSlvNod),
5 fNormal(nContact), shape(nFacNod,nContact),
6 coordSlv(nDir,nSlvNod), coordMst(nDir,nMstNod),
7 dirCosSl(nDir,nContact), dircosN(nDir,nContact),
8 jSlvUid(nSlvNod), jMstUid(nMstNod),
9 jConSlvid(nContact), jConMstid(nFacNod,nContact)
1 tempSlv(nContact), preDefSlv(nContact,nPred),
2 tempMst(numDefTfv), preDefMst(numDefTfv,nPred)

  user coding to define fTangential
  and, optionally, statev

  return
end

```

---

**Variable to be defined****fTangential (nFricDir, nContact)**

This array must be updated to the current values of the frictional force components for all contact points in the local tangent directions. See Figure 1.2.4–1 and Figure 1.2.4–2 for definition of the local coordinate system. This array will be zero (no friction force) until you reset it.

---

**Variable that can be updated****statev (nstateVar, nSlvNod)**

This array contains the user-defined solution-dependent state variables for all the nodes on the slave surface. You define the size of this array (see “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual, for more information). This array will be passed in containing the values of these variables prior to the call to user subroutine **VFRIC**. If any of the solution-dependent state variables is being used in conjunction with the friction behavior, it must be updated in this subroutine.

The state variables are available even for slave nodes that are not in contact. This may be useful when, for example, the state variables need to be reset for slave nodes that are not in contact.

### **Variables passed in for information**

---

#### **kStep**

Step number.

#### **kInc**

Increment number.

#### **nContact**

Number of contacting slave nodes.

#### **nFacNod**

Number of nodes on each master surface facet (**nFacNod** is 2 for two-dimensional surfaces, **nFacNod** is 4 for three-dimensional surfaces). If the master surface is an analytical rigid surface, this variable is passed in as 0.

#### **nSlvNod**

Number of slave nodes.

#### **nMstNod**

Number of master surface nodes, if the master surface is made up of facets. If the master surface is an analytical rigid surface, this variable is passed in as 0.

#### **nFricDir**

Number of tangent directions at the contact points (**nFricDir = nDir - 1**).

#### **nDir**

Number of coordinate directions at the contact points. (In a three-dimensional model **nDir** will be two if the surfaces in the contact pair are two-dimensional analytical rigid surfaces or are formed by two-dimensional elements.)

#### **nStateVar**

Number of user-defined state variables.

#### **nProps**

User-specified number of property values associated with this friction model.

#### **nTemp**

1 if the temperature is defined and 0 if the temperature is not defined.

#### **nPred**

Number of predefined field variables.

**numDefTfv**

Equal to **nContact** if the master surface is made up of facets. If the master surface is an analytical rigid surface, this variable is passed in as 1.

**jSlvUid(nSlvNod)**

This array lists the user-defined global node numbers (or internal node numbers for models defined in terms of an assembly of part instances) of the nodes on the slave surface.

**jMstUid(nMstNod)**

This array lists the user-defined global node numbers (or internal node numbers for models defined in terms of an assembly of part instances) of the nodes on the master surface. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**jConSlvid(nContact)**

This array lists the surface node numbers of the slave surface nodes that are in contact.

**jConMstid(nFacNod, nContact)**

This array lists the surface node numbers of the master surface nodes that make up the facet with which each contact point is in contact. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**timStep**

Value of step time.

**timGlb**

Value of total time.

**dtimCur**

Current increment in time from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ .

**surfInt**

User-specified surface interaction name, left justified.

**surfSlv**

Slave surface name.

**surfMst**

Master surface name.

**lContType**

Contact type flag. This flag is set based on the type of constraint enforcement method (see “Contact constraint enforcement methods in Abaqus/Explicit,” Section 34.2.3 of the Abaqus Analysis User’s Manual) being used: 1 for kinematic contact and 2 for penalty contact. Stick conditions are satisfied exactly with the kinematic contact algorithm; they are satisfied only approximately (subject to an automatically chosen penalty stiffness value) with the penalty contact algorithm.

**dSlipFric (nDir, nContact)**

This array contains the incremental frictional slip during the current time increment for each contact point in the current local coordinate system. These incremental slips correspond to tangential motion in the time increment from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ . For penalty contact this incremental slip is used to define the local coordinate system at each contact point (see Figure 1.2.4–1 and Figure 1.2.4–2) so that only the first component of **dSlipFric** can be nonzero in the local system. The contact points for kinematic contact are determined based on penetrations detected in the predicted configuration (at  $t = t_{curr} + \Delta t$ ), and the predicted incremental slip direction is used to define the local coordinate system at each contact point. If the slip direction changes between increments, **dSlipFric** may have a nonzero component in the local 2-direction and, if the surface is faceted and the contact point moves from one facet to another, in the local 3-direction.

**fStickForce (nContact)**

This array contains the magnitude of frictional force required to enforce stick conditions at each contact point. For kinematic contact this force corresponds to no slip; for penalty contact this force depends on the previous frictional force, the value of the penalty stiffness, and the previous incremental slip. The penalty stiffness is assigned automatically. Occasionally, during recovery of elastic slip associated with the penalty method, the stick force will be assigned a negative value.

**fTangPrev (nDir, nContact)**

This array contains the values of the frictional force components calculated in the previous increment but provided in the current local coordinate system (zero for nodes that were not in contact).

**fNormal (nContact)**

This array contains the magnitude of the normal force for the contact points applied at the end of current time increment; i.e., at time  $t = t_{curr}$ .

**frictionWork**

This variable contains the value of the total frictional dissipation in the entire model from the beginning of the analysis. The units are energy per unit area.

**shape (nFacNod, nContact)**

For each contact point this array contains the shape functions of the nodes of its master surface facet, evaluated at the location of the contact point. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**coordSlv (nDir, nSlvNod)**

Array containing the **nDir** components of the current coordinates of the slave nodes.

**coordMst (nDir, nMstNod)**

Array containing the **nDir** components of the current coordinates of the master nodes. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**dirCosSl(nDir, nContact)**

Direction cosines of the incremental slip at the contact points.

**dircosN(nDir, nContact)**

Direction cosines of the normals to the master surface at the contact points.

**props(nProps)**

User-specified vector of property values to define the frictional behavior between the contacting surfaces.

**areaSlv(nSlvNod)**

Area associated with the slave nodes (equal to 1 for node-based surface nodes).

**tempSlv(nContact)**

Current temperature at the slave nodes.

**preDefSlv(nContact, nPred)**

Current user-specified predefined field variables at the slave nodes (initial values at the beginning of the analysis and current values during the analysis).

**tempMst(numDefTfv)**

Current temperature at the nearest points on the master surface.

**preDefMst(numDefTfv, nPred)**

Current user-specified predefined field variables at the nearest points on the master surface (initial values at the beginning of the analysis and current values during the analysis).

## 1.2.5 VFRIC\_COEF: User subroutine to define the frictional coefficient for contact surfaces.

**Product:** Abaqus/Explicit

### References

---

- “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual
- \*FRICTION
- “**VFRIC**, **VFRIC\_COEF**, and **VFRICTION**,” Section 4.1.30 of the Abaqus Verification Manual

### Overview

---

User subroutine **VFRIC\_COEF**:

- can be used to define the isotropic frictional coefficient between contacting surfaces;
- corresponds to the classical Coulomb friction model; and
- can be used only with the general contact algorithm.

### User subroutine interface

---

```

subroutine vfric_coef (
C Write only -
      *   fCoef, fCoefDeriv,
C Read only -
      *   nBlock, nProps, nTemp, nFields,
      *   jFlags, rData,
      *   surfInt, surfSlv, surfMst,
      *   props, slipRate, pressure,
      *   tempAvg, fieldAvg )
C
      include 'vaba_param.inc'
C
      dimension fCoef(nBlock),
      *   fCoefDeriv(nBlock,3),
      *   props(nProps),
      *   slipRate(nBlock),
      *   pressure(nBlock),
      *   tempAvg(nBlock),
      *   fieldAvg(nBlock,nFields)
C
      parameter( iKStep    = 1,

```

## VFRIC\_COEF

```
*          iKInc    = 2,
*          nFlags   = 2 )
C
parameter( iTimStep = 1,
*          iTimGlb  = 2,
*          iDTimCur = 3,
*          nData    = 3 )
C
dimension jFlags(nFlags), rData(nData)
C
character*80 surfInt, surfSlv, surfMst
C
user coding to define fCoef
C
return
end
```

### Variables to be defined

---

#### **fCoef(nBlock)**

This array must be updated to the current values of the friction coefficient for all contacting points.

#### **fCoefDeriv(nBlock,3)**

This array is not applicable to Abaqus/Explicit analyses.

### Variables passed in for information

---

#### **nBlock**

Number of contacting points to be processed in this call to **VFRIC\_COEF**.

#### **nProps**

User-specified number of property values associated with this friction model.

#### **nTemp**

1 if the temperature is defined and 0 if the temperature is not defined.

#### **nFields**

Number of user-specified field variables.

#### **jFlag(1)**

Step number.

#### **jFlag(2)**

Increment number.

**rData(1)**

Value of step time.

**rData(2)**

Value of total time.

**rData(3)**

Current increment in time from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ .

**surfInt**

User-specified surface interaction name, left justified.

**surfSlv**

Slave surface name, not applicable to general contact.

**surfMst**

Master surface name, not applicable to general contact.

**props(nProps)**

User-specified vector of property values to define the frictional coefficient at contacting points.

**slipRate(nBlock)**

This array contains the rate of tangential slip at the contacting points for the current time increment.

**pressure(nBlock)**

This array contains the pressure at the contacting points applied at the end of the current time increment.

**tempSlv(nBlock)**

Average current temperature between the master and slave surfaces at the contacting points.

**fieldSlv(nFields,nBlock)**

Average current value of all the user-specified field variables between the master and slave surfaces at the contacting points.



## 1.2.6 VFRICITION: User subroutine to define frictional behavior for contact surfaces.

**Product:** Abaqus/Explicit

### References

---

- “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual
- \*FRICTION
- “**VFRIC**, **VFRIC\_COEF**, and **VFRICITION**,” Section 4.1.30 of the Abaqus Verification Manual

### Overview

---

User subroutine **VFRICITION**:

- can be used to define the frictional behavior between contacting surfaces;
- can be used when the classical Coulomb friction model is too restrictive and a more complex definition of shear transmission between contacting surfaces is required;
- must provide the entire definition of shear interaction between the contacting surfaces;
- can use and update solution-dependent state variables for node-to-face and node-to-analytical rigid surface contact;
- cannot be used in conjunction with softened tangential surface behavior; and
- can be used only with the general contact algorithm.

### Terminology

---

The use of user subroutine **VFRICITION** requires familiarity with the following terminology.

#### Surface node numbers

The “surface node number” refers to the position of a particular node in the list of nodes on the surface. For example, there are **nSlvNod** nodes on the slave surface. Number  $n, n = 1, 2, \dots, \text{nSlvNod}$ , is the surface node number of the  $n$ th node in this list; **jSlvUid**( $n$ ) is the user-defined global number of this node. An Abaqus/Explicit model can be defined in terms of an assembly of part instances (see “Defining an assembly,” Section 2.9.1 of the Abaqus Analysis User’s Manual). In such models a node number in **jSlvUid** is an internally generated node number. If the original node number and part instance name are required, call the utility routine **VGETPARTINFO** (see “Obtaining part information,” Section 2.1.5).

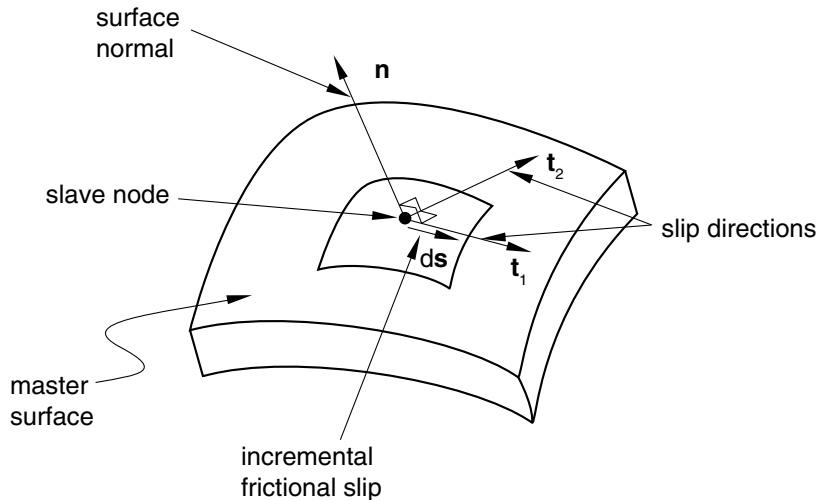
#### Contact points

The nodes on the slave surface that are in contact in the current time increment are defined as “contact points.” The number of contact points is passed into user subroutine **VFRICITION** as **nContact**. The array **jConSlvid**(**nContact**) gives the surface node numbers for the contact points.

## Local coordinate system

---

A local coordinate system is defined for each contact point to facilitate specification of frictional forces and incremental slip. The local 1-direction is tangential to the master surface; it is defined by  $t_1 = ds/|ds|$ , where  $ds$  is the incremental slip vector. The incremental slip vector used to define  $t_1$  corresponds to the incremental slip in the current time increment. The master surface normal direction,  $n$ , is the local 3-direction. The local 2-direction is given by  $t_2 = n \times t_1$ , which is also tangent to the master surface. The vectors are shown in Figure 1.2.6–1. The direction cosines for  $t_1$  and  $n$  with respect to the global coordinate system are available in **dirCosS1** and **dirCosN**, respectively. In the case of zero incremental slip ( $|ds| = 0$ ) we choose an arbitrary direction for  $t_1$  that is orthogonal to the normal direction,  $n$ .



**Figure 1.2.6–1** Local coordinate system for three-dimensional contact with **VFRICTION**.

## Frictional forces

---

You specify the frictional force, **fTangential**, at each contact point in local coordinates in this subroutine. The array **fTangential** is dimensioned such that only the tangential components can be specified. Any components of the frictional force that are not specified will remain equal to zero. For isotropic friction, only the first component of the frictional force need be specified since the second component should be zero. A “stick force” at each contact point is provided in the array **fStickForce** to assist you in setting appropriate frictional force values. The stick force is the force required to prevent additional “plastic” slipping. The stick force at each contact point is provided as a scalar value as it would act in the direction opposite to  $t_1$ . The stick force is computed prior to calling user subroutine **VFRICTION**. The first component of the frictional force should be in the range between zero and the

negative of the stick force value. Typically, the stick force will be positive and the first component of the applied frictional force will be negative, opposing the incremental slip. Penalty contact includes an elastic slip regime due to finite penalty stiffness; so occasionally the stick force will be negative during recovery of elastic slip, indicating that it is appropriate for the first component of the frictional force to be positive (i.e., acting in the same direction as the incremental slip). A noisy or unstable solution is likely to result if the first component of **fTangential** is set outside the range between zero and the negative of the stick force value.

After user subroutine **VFRICITION** is called, frictional forces that oppose the forces specified at the contact points are distributed to the master nodes.

## User subroutine interface

---

```

        subroutine vfriction (
C Write only -
        *   fTangential,
C Read/Write -
        *   state,
C Read only -
        *   nBlock, nBlockAnal, nBlockEdge,
        *   nNodState, nNodSlv, nNodMst,
        *   nFricDir, nDir,
        *   nStates, nProps, nTemp, nFields,
        *   jFlags, rData,
        *   surfInt, surfSlv, surfMst,
        *   jConSlvUid, jConMstUid, props,
        *   dSlipFric, fStickForce, fTangPrev, fNormal,
        *   areaSlv, dircosN, dircosSlv,
        *   shapeSlv, shapeMst,
        *   coordSlv, coordMst,
        *   velSlv, velMst,
        *   tempSlv, tempMst,
        *   fieldSlv, fieldMst )
C
        include `vaba_param.inc'
C
        dimension fTangential(nFricDir,nBlock),
        *   state(nStates,nNodState,nBlock),
        *   jConSlvUid(nNodSlv,nBlock),
        *   jConMstUid(nNodMst,nBlockAnal),
        *   props(nProps),
        *   dSlipFric(nDir,nBlock),
        *   fStickForce(nBlock),

```

```

*   fTangPrev(nDir,nBlock) ,
*   fNormal(nBlock) ,
*   areaSlv(nBlock) ,
*   dircosN(nDir,nBlock) ,
*   dircosS1(nDir,nBlock) ,
*   shapeSlv(nNodSlv,nBlockEdge) ,
*   shapeMst(nNodMst,nBlockAnal) ,
*   coordSlv(nDir,nNodSlv,nBlock) ,
*   coordMst(nDir,nNodMst,nBlockAnal) ,
*   velSlv(nDir,nNodSlv,nBlock) ,
*   velMst(nDir,nNodMst,nBlockAnal) ,
*   tempSlv(nBlock) ,
*   tempMst(nBlockAnal) ,
*   fieldSlv(nFields,nBlock) ,
*   fieldMst(nFields,nBlockAnal)

C
    parameter( iKStep      = 1,
*              iKInc       = 2,
*              iLConType   = 3,
*              nFlags      = 3 )

C
    parameter( iTimStep     = 1,
*              iTimGlb      = 2,
*              iDTimCur     = 3,
*              iFrictionWork = 4,
*              nData        = 4 )

C
    dimension jFlags(nFlags), rData(nData)

C
    character*80 surfInt, surfSlv, surfMst

C
    user coding to define fTangential
    and, optionally, state

C
    return
end

```

**Variable to be defined****fTangential(nFricDir,nBlock)**

This array must be updated to the current values of the frictional force components for all contact points in the local tangent directions. See Figure 1.2.6–1 for a definition of the local coordinate system. This array will be zero (no friction force) until it is set.

## Variable that can be updated

---

### **state (nStates, nNodState, nBlock)**

This array contains the user-defined, solution-dependent state variables for all the nodes on the slave surface. The use of state variables is applicable for node-to-face and node-to-analytical rigid surface contact. See “Frictional behavior,” Section 33.1.5 of the Abaqus Analysis User’s Manual, for more information on the size of this array. This array will be passed in containing the values of these variables prior to the call to user subroutine **VFRICITION**.

If any of the solution-dependent state variables are being used in conjunction with the friction behavior, they must be updated in this subroutine. These state variables need to be updated with care, as a slave node can be in contact with multiple master surfaces. Such a slave node may be passed into the user subroutine at a given increment multiple times, possibly on separate calls to the user subroutine, and you may end up advancing the state variables for that node multiple times for a single time increment. One trick to keep track of whether or not a node state is advanced is to use one of the state variables exclusively for this purpose. You could set that selected state variable to the current increment number and update the state only if it is not already set to the current increment number.

## Variables passed in for information

---

### **nBlock**

Number of contacting points to be processed in this call to **VFRICITION**.

### **nBlockAnal**

1 for analytical rigid master surface; **nBlock** otherwise.

### **nBlockEdge**

**nBlock** for edge-type slave surface; 1 otherwise.

### **nNodState**

1 for node-to-face contact and node-to-analytical rigid surface contact.

### **nNodSlv**

1 for node-to-face and node-to-analytical rigid surface contact; 2 for edge-to-edge contact.

### **nNodMst**

1 for analytical rigid master surface; 2 for edge-type master surface; 4 for facet-type master surface.

### **nFricDir**

Number of tangent directions at the contact points (**nFricDir = nDir - 1**).

### **nDir**

Number of coordinate directions at the contact points (equal to 3).

### **nStates**

Number of user-defined state variables.

## VFRICITION

### **nProps**

User-specified number of property values associated with this friction model.

### **nTemp**

1 if the temperature is defined and 0 if the temperature is not defined.

### **nFields**

Number of predefined field variables.

### **jFlag(1)**

Step number.

### **jFlag(2)**

Increment number.

### **jFlag(3)**

1 for node-to-face contact, 2 for edge-to-edge contact, and 3 for node-to-analytical rigid surface contact.

### **rData(1)**

Value of step time.

### **rData(2)**

Value of total time.

### **rData(3)**

Current increment in time from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ .

### **rData(4)**

This variable contains the value of the total frictional dissipation in the entire model from the beginning of the analysis. The units are energy per unit area.

### **surfInt**

User-specified surface interaction name, left justified.

### **surfSlv**

Slave surface name, not applicable to general contact.

### **surfMst**

Master surface name, not applicable to general contact.

### **jConSlvUid(nNodSlv,nBlock)**

This array lists the surface node numbers of the slave surface nodes that are in contact.

### **jConMstUid(nNodMst,nBlockAnal)**

This array lists the surface node numbers of the master surface nodes that make up the facet with which each slave node is in contact.

**props (nProps)**

User-specified vector of property values to define the frictional behavior between the contacting surfaces.

**dSlipFric (nDir, nBlock)**

This array contains the incremental frictional slip during the current time increment for each contacting point in the current local coordinate system. These incremental slips correspond to tangential motion in the time increment from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ . This incremental slip is used to define the local coordinate system at each contact point (see Figure 1.2.6-1) so that only the first component of **dSlipFric** can be nonzero in the local system. If the slip direction changes between increments, **dSlipFric** may have a nonzero component in the local 2-direction and, if the surface is faceted and the contact point moves from one facet to another, in the local 3-direction.

**fStickForce (nBlock)**

This array contains the magnitude of frictional force required to enforce stick conditions at each contact point. This force depends on the previous frictional force, the value of the penalty stiffness, and the previous incremental slip. The penalty stiffness is assigned automatically. Occasionally, during recovery of elastic slip associated with the penalty method, the stick force will be assigned a negative value.

**fTangPrev (nDir, nBlock)**

This array contains the values of the frictional force components calculated in the previous increment but provided in the current local coordinate system (zero for nodes that were not in contact).

**fNormal (nBlock)**

This array contains the magnitude of the normal force for the contact points applied at the end of current time increment; i.e., at time  $t = t_{curr}$ .

**areaSlv (nBlock)**

Area associated with the slave nodes (equal to 1 for node-based surface nodes).

**dircosN (nDir, nBlock)**

Direction cosines of the normals to the master surface at the contact points.

**dirCosS1 (nDir, nBlock)**

Direction cosines of the incremental slip at the contact points. The direction cosines are undefined (all components zero) if the incremental frictional slip is zero.

**shapeSlv (nNodSlv, nBlockEdge)**

For each contact point this array contains the shape functions of the nodes of its slave surface, evaluated at the location of the contact point. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**shapeMst (nNodMst, nBlockAnal)**

For each contact point this array contains the shape functions of the nodes of its master surface, evaluated at the location of the contact point. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**coordSlv (nDir, nNodSlv, nBlock)**

Array containing the **nDir** components of the current coordinates of the slave nodes.

**coordMst (nDir, nNodMst, nBlockAnal)**

Array containing the **nDir** components of the current coordinates of the master nodes. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**velSlv (nDir, nNodSlv, nBlock)**

Array containing the **nDir** components of the current velocity of the slave nodes.

**velMst (nDir, nNodMst, nBlockAnal)**

Array containing the **nDir** components of the current velocity of the master nodes. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**tempSlv (nBlock)**

Current temperature at the slave nodes.

**tempMst (nBlockAnal)**

Current temperature at the nearest points on the master surface.

**fieldSlv (nFields, nBlock)**

Current user-specified predefined field variables at the slave nodes (initial values at the beginning of the analysis and current values during the analysis).

**fieldMst (nFields, nBlockAnal)**

Current user-specified predefined field variables at the nearest points on the master surface (initial values at the beginning of the analysis and current values during the analysis).

## 1.2.7 VUAMP: User subroutine to specify amplitudes.

**Product:** Abaqus/Explicit

### References

---

- “Amplitude curves,” Section 30.1.2 of the Abaqus Analysis User’s Manual
- \*AMPLITUDE
- \*OUTPUT

### Overview

---

User subroutine **VUAMP**:

- allows you to define the current value of an amplitude definition as a function of time;
- can be used to model control engineering aspects of your system when sensors are used (sensor values are from the beginning of the increment);
- can use a predefined number of state variables in its definition; and
- can optionally compute the derivatives and integrals of the amplitude function.

### Explicit solution dependence

---

The solution dependence introduced in this user subroutine is explicit: all data passed in the subroutine for information or to be updated are values at the beginning of that increment.

### User subroutine interface

---

```

SUBROUTINE VUAMP(
*      ampName, time, ampValueOld, dt, nSvars, svars,
*      lFlagsInfo,
*      nSensor, sensorValues, sensorNames, jSensorLookUpTable,
*      AmpValueNew,
*      lFlagsDefine,
*      AmpDerivative, AmpSecDerivative, AmpIncIntegral)
C
INCLUDE 'VABA_PARAM.INC'

C      time indices
parameter (iStepTime      = 1,
*              iTotalTime     = 2,
*              nTime          = 2)
C      flags passed in for information

```

```

        parameter (iInitialization = 1,
*                  iRegularInc   = 2,
*                  ikStep        = 3,
*                  nFlagsInfo    = 3)
C      optional flags to be defined
        parameter (iComputeDeriv   = 1,
*                  iComputeSecDeriv = 2,
*                  iComputeInteg   = 3,
*                  iStopAnalysis   = 4,
*                  iConcludeStep   = 5,
*                  nFlagsDefine    = 5)
        dimension time(nTime), lFlagsInfo(nFlagsInfo),
*                  lFlagsDefine(nFlagsDefine),
*                  sensorValues(nSensor),
*                  sVars(nSvars)

        character*80 sensorNames(nSensor)
        character*80 ampName
        dimension jSensorLookUpTable(*)

user coding to define AmpValueNew, and
optionally lFlagsDefine, AmpDerivative, AmpSecDerivative,
AmpIncIntegral

        RETURN
        END

```

**Variable to be defined**

---

**AmpValueNew**

Current value of the amplitude.

**Variables that can be updated**

---

**lFlagsDefine**

Integer flag array to determine whether the computation of additional quantities is necessary or to set step continuation requirements.

**lFlagsDefine(iComputeDeriv)**

If set to 1, you must provide the computation of the amplitude derivative. The default is 0, which means that Abaqus computes the derivative automatically.

**lFlagsDefine(iComputeSecDeriv)**

If set to 1, you must provide the computation of the amplitude second derivative. The default is 0, which means that Abaqus computes the second derivative automatically.

**lFlagsDefine(iComputeInteg)**

If set to 1, you must provide the computation of the amplitude incremental integral. The default is 0, which means that Abaqus computes the incremental integral automatically.

**lFlagsDefine(iStopAnalysis)**

If set to 1, the analysis will be stopped and an error message will be issued. The default is 0, which means that Abaqus will not stop the analysis.

**lFlagsDefine(iConcludeStep)**

If set to 1, Abaqus will conclude the step execution and advance to the next step (if a next step is available). The default is 0.

**svars**

An array containing the values of the solution-dependent state variables associated with this amplitude definition. The number of such variables is **nsvars** (see above). You define the meaning of these variables.

This array is passed into **VUAMP** containing the values of these variables at the start of the current increment. In most cases they should be updated to be the values at the end of the increment.

**AmpDerivative**

Current value of the amplitude derivative.

**AmpSecDerivative**

Current value of the amplitude second derivative.

**AmpIncIntegral**

Current value of the amplitude incremental integral.

**Variables passed in for information**

---

**ampName**

User-specified amplitude name, left justified.

**time(iStepTime)**

Current value of step time.

**time(iTotalTime)**

Current value of total time.

**ampValueOld**

Old value of the amplitude from the previous increment.

**dt**

Current stable time increment.

**nSvars**

User-defined number of solution-dependent state variables associated with this amplitude definition.

**lFlagsInfo**

Integer flag array with information regarding the current call to **VUAMP**:

**lFlagsInfo(iInitialization)**

This flag is equal to 1 if **VUAMP** is called from the initialization phase of each step and is set to 0 otherwise.

**lFlagsInfo(iRegularInc)**

This flag is equal to 1 if **VUAMP** is called from a regular increment and is set to 0 otherwise.

**lFlagsInfo(ikStep)**

Step number.

**nSensor**

Total number of sensors in the model.

**sensorValues**

Array with sensor values at the end of the previous increment. Each sensor value corresponds to a history output variable associated with the output database request defining the sensor.

**sensorNames**

Array with user-defined sensor names in the entire model, left justified. Each sensor name corresponds to a sensor value provided with the output database request. All names will be converted to uppercase characters if lowercase or mixed-case characters were used in their definition.

**jSensorLookUpTable**

Variable that must be passed into the utility functions **IVGETSENSORID** and **VGETSENSORVALUE**.

**Example: Amplitude definition using sensor and state variables**

```

C      user amplitude subroutine
      Subroutine VUAMP(
C      passed in for information and state variables
*      ampName, time, ampValueOld, dt, nSvars, svars,
*      lFlagsInfo,
*      nSensor, sensorValues, sensorNames,
*      jSensorLookUpTable,
C      to be defined
*      ampValueNew,
```

```

*      lFlagsDefine,
*      AmpDerivative, AmpSecDerivative, AmpIncIntegral)

include 'vaba_param.inc'

C   svars - additional state variables, similar to (V)UEL
dimension sensorValues(nSensor), svars(nSvars)
character*80 sensorNames(nSensor)
character*80 ampName

C   time indices
parameter( iStepTime      = 1,
*           iTotTime       = 2,
*           nTime          = 2)
C   flags passed in for information
parameter( iInitialization = 1,
*           iRegularInc   = 2,
*           ikStep         = 3,
*           nFlagsInfo    = 3)
C   optional flags to be defined
parameter( iComputeDeriv  = 1,
*           iComputeSecDeriv= 2,
*           iComputeInteg = 3,
*           iStopAnalysis = 4,
*           iConcludeStep = 5,
*           nFlagsDefine  = 5)

parameter( tStep=0.18, tAccelerateMotor = .00375,
*           omegaFinal=23.26)

dimension time(nTime), lFlagsInfo(nFlagsInfo),
*           lFlagsDefine(nFlagsDefine)
dimension jSensorLookUpTable(*)

lFlagsDefine(iComputeDeriv)    = 1
lFlagsDefine(iComputeSecDeriv) = 1

c   get sensor value
vTrans_CU1  = vGetSensorValue('HORIZ_TRANSL_MOTION',
*                               jSensorLookUpTable,
*                               sensorValues)

```

```
if (ampName(1:22) .eq. 'MOTOR_WITH_STOP_SENSOR' ) then
    if (lFlagsInfo(iInitialization).eq.1) then
        ampValueNew      = ampValueOld

        svars(1) = 0.0
        svars(2) = 0.0
    else
        tim = time(iStepTime)

c      ramp up the angular rot velocity  of the electric
c      motor after which hold constant
        if (tim .le. tAccelerateMotor) then
            ampValueNew = omegaFinal*tim/tAccelerateMotor

        else
            ampValueNew = omegaFinal
        end if

c      retrieve old sensor value
        vTrans_CU1_old = svars(1)

c      detect a zero crossing and count the number of
c      crossings
        if (vTrans_CU1_old*vTrans_CU1 .le. 0.0 .and.
*             tim .gt. tAccelerateMotor ) then
            svars(2) = svars(2) + 1.0
        end if
        nrCrossings = int(svars(2))

c      stop the motor if sensor crosses zero the second
c      time
        if (nrCrossings.eq.2) then
            ampValueNew = 0.0
            lFlagsDefine(iConcludeStep)=1
        end if

c      store sensor value
        svars(1) = vTrans_CU1
```

```
    end if
end if
```

```
return
end
```



## 1.2.8 VUANISOHYPER\_INV: User subroutine to define anisotropic hyperelastic material behavior using the invariant formulation.

**Product:** Abaqus/Explicit

### References

---

- “Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual
- \*ANISOTROPIC HYPERELASTIC
- “**VUANISOHYPER\_INV** and **VUANISOHYPER\_INV**,” Section 4.1.13 of the Abaqus Verification Manual

### Overview

---

User subroutine **VUANISOHYPER\_INV**:

- can be used to define the strain energy potential of anisotropic hyperelastic materials as a function of an irreducible set of scalar invariants;
- will be called for blocks of material calculation points for which the material definition contains user-defined anisotropic hyperelastic behavior with invariant-based formulation (“Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual);
- can use and update solution-dependent state variables;
- can use any field variables that are passed in; and
- requires that the values of the derivatives of the strain energy density function be defined with respect to the scalar invariants.

### Enumeration of invariants

---

To facilitate coding and provide easy access to the array of invariants passed to user subroutine **VUANISOHYPER\_INV**, an enumerated representation of each invariant is introduced. Any scalar invariant can, therefore, be represented uniquely by an *enumerated* invariant,  $I_n^*$ , where the subscript  $n$  denotes the order of the invariant according to the enumeration scheme in the following table:

Invariant	Enumeration, $n$
$\bar{I}_1$	1
$\bar{I}_2$	2
$J$	3
$\bar{I}_{4(\alpha\beta)}$	$4 + 2(\alpha - 1) + \beta(\beta - 1); \quad \alpha \leq \beta$
$\bar{I}_{5(\alpha\beta)}$	$5 + 2(\alpha - 1) + \beta(\beta - 1); \quad \alpha \leq \beta$

For example, in the case of three families of fibers there are a total of 15 invariants:  $\bar{I}_1$ ,  $\bar{I}_2$ ,  $J$ , six invariants of type  $\bar{I}_{4(\alpha\beta)}$ , and six invariants of type  $\bar{I}_{5(\alpha\beta)}$ , with  $\alpha, \beta = 1, 2, 3$  ( $\alpha \leq \beta$ ). The following correspondence exists between each of these invariants and their enumerated counterpart:

Enumerated invariant	Invariant
$I_1^*$	$\bar{I}_1$
$I_2^*$	$\bar{I}_2$
$I_3^*$	$J$
$I_4^*$	$\bar{I}_{4(11)}$
$I_5^*$	$\bar{I}_{5(11)}$
$I_6^*$	$\bar{I}_{4(12)}$
$I_7^*$	$\bar{I}_{5(12)}$
$I_8^*$	$\bar{I}_{4(22)}$
$I_9^*$	$\bar{I}_{5(22)}$
$I_{10}^*$	$\bar{I}_{4(13)}$
$I_{11}^*$	$\bar{I}_{5(13)}$
$I_{12}^*$	$\bar{I}_{4(23)}$
$I_{13}^*$	$\bar{I}_{5(23)}$
$I_{14}^*$	$\bar{I}_{4(33)}$
$I_{15}^*$	$\bar{I}_{5(33)}$

A similar scheme is used for the array **zeta** of terms  $\zeta_{\alpha\beta} = \mathbf{A}_\alpha \cdot \mathbf{A}_\beta$ . Each term can be represented uniquely by an enumerated counterpart  $\zeta_m^*$ , as shown below:

Dot product	Enumeration, $m$
$\zeta_{\alpha\beta}$	$\alpha + \frac{1}{2}(\beta - 2)(\beta - 1); \quad \alpha < \beta$

As an example, for the case of three families of fibers there are three  $\zeta_{\alpha\beta}$  terms:  $\zeta_{12}$ ,  $\zeta_{13}$ , and  $\zeta_{23}$ . These are stored in the **zeta** array as  $(\zeta_1^*, \zeta_2^*, \zeta_3^*)$ .

---

## Storage of arrays of derivatives of energy function

The components of the array **duDi** of first derivatives of the strain energy potential with respect to the scalar invariants,  $\partial U / \partial I_i^*$ , are stored using the enumeration scheme discussed above for the scalar invariants.

The elements of the array **d2uDiDi** of second derivatives of the strain energy function,  $\partial^2 U / \partial I_i^* \partial I_j^*$ , are laid out in memory using triangular storage: if  $k$  denotes the component in this array corresponding to the term  $\partial^2 U / \partial I_i^* \partial I_j^*$ , then  $k = i + j \times (j - 1)/2$ ; ( $i \leq j$ ). For example, the term  $\partial^2 U / \partial I_2^* \partial I_5^*$  is stored in component  $k = 2 + (5 \times 4)/2 = 12$  in the **d2uDiDi** array.

---

## Special considerations for shell elements

When **VUANISOHYPER\_INV** is used to define the material response of shell elements, Abaqus/Explicit cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element's transverse shear stiffness. See "Shell section behavior," Section 26.6.4 of the Abaqus Analysis User's Manual, for guidelines on choosing this stiffness.

---

## Material point deletion

Material points that satisfy a user-defined failure criterion can be deleted from the model (see "User-defined mechanical material behavior," Section 23.8.1 of the Abaqus Analysis User's Manual). You must specify the state variable number controlling the element deletion flag when you allocate space for the solution-dependent state variables, as explained in "User-defined mechanical material behavior," Section 23.8.1 of the Abaqus Analysis User's Manual. The deletion state variable should be set to a value of one or zero in **VUANISOHYPER\_INV**. A value of one indicates that the material point is active, and a value of zero indicates that Abaqus/Explicit should delete the material point from the model by setting the stresses to zero. The structure of the block of material points passed to user subroutine **VUANISOHYPER\_INV** remains unchanged during the analysis; deleted material points are not removed from the block. Abaqus/Explicit will "freeze" the values of the invariants passed to **VUANISOHYPER\_INV** for all deleted material points; that is, the values remain constant after deletion is triggered. Once a material point has been flagged as deleted, it cannot be reactivated.

---

## User subroutine interface

```

subroutine vuanisohyper_inv (
C Read only (unmodifiable) variables -
  1      nblock, nFiber, nInv,
  2      jElem, kIntPt, kLayer, kSecPt,
  3      cmname,
  4      nstatev, nfieldv, nprops,
  5      props, tempOld, tempNew, fieldOld, fieldNew,
  6      stateOld, sInvariant, zeta,
C Write only (modifiable) variables -

```

```

7      uDev, duDi, d2uDiDi,
8      stateNew )
C
      include 'vaba_param.inc'
C
      dimension props(nprops),
1  tempOld(nblock),
2  fieldOld(nblock,nfieldv),
3  stateOld(nblock,nstatev),
4  tempNew(nblock),
5  fieldNew(nblock,nfieldv),
6  sInvariant(nblock,nInv),
7  zeta(nblock,nFiber*(nFiber-1)/2),
8  uDev(nblock), duDi(nblock,nInv),
9  d2uDiDi(nblock,nInv*(nInv+1)/2),
* stateNew(nblock,nstatev)
C
      character*80 cmname
C

do 100 km = 1,nblock
  user coding
100 continue

      return
      end

```

---

**Variables to be defined****udev (nblock)**

$\tilde{U}_{dev}$ , the deviatoric part of the strain energy density of the primary material response. This quantity is needed only if the current material definition also includes Mullins effect (see “Mullins effect,” Section 19.6.1 of the Abaqus Analysis User’s Manual).

**duDi (nblock, nInv)**

Array of derivatives of strain energy potential with respect to the scalar invariants,  $\partial U / \partial I_i^*$ , ordered using the enumeration scheme discussed above.

**d2uDiDi (nblock, nInv\*(nInv+1)/2)**

Arrays of second derivatives of strain energy potential with respect to the scalar invariants (using triangular storage),  $\partial^2 U / \partial I_i^* \partial I_j^*$ .

**stateNew(nblock,nstatev)**

State variables at each material point at the end of the increment. You define the size of this array by allocating space for it (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information).

**Variables passed in for information**

---

**nblock**

Number of material points to be processed in this call to **VUANISOHYPER\_STRAIN**.

**nFiber**

Number of families of fibers defined for this material.

**nInv**

Number of scalar invariants.

**jElem(nblock)**

Array of element numbers.

**kIntPt**

Integration point number.

**kLayer**

Layer number (for composite shells).

**kSecPt**

Section point number within the current layer.

**cmname**

User-specified material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, you should not use “ABQ\_” as the leading string for **cmname**.

**nstatev**

Number of user-defined state variables that are associated with this material type (you define this as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**props(nprops)**

User-supplied material properties.

## VUANISOHYPER\_INV

### **tempOld(nblock)**

Temperatures at each material point at the beginning of the increment.

### **tempNew(nblock)**

Temperatures at each material point at the end of the increment.

### **fieldOld(nblock, nfieldv)**

Values of the user-defined field variables at each material point at the beginning of the increment.

### **fieldNew(nblock, nfieldv)**

Values of the user-defined field variables at each material point at the end of the increment.

### **stateOld(nblock, nstatev)**

State variables at each material point at the beginning of the increment.

### **sInvariant(nblock, nInv)**

Array of scalar invariants,  $I_i^*$ , at each material point at the end of the increment. The invariants are ordered using the enumeration scheme discussed above.

### **zeta(nblock, nFiber\*(nFiber-1)/2) )**

Array of dot product between the directions of different families of fiber in the reference configuration,  $\zeta_{\alpha\beta} = \mathbf{A}_\alpha \cdot \mathbf{A}_\beta$ . The array contains the enumerated values  $\zeta_m^*$  using the scheme discussed above.

---

### **Example: Using more than one user-defined anisotropic hyperelastic material model**

To use more than one user-defined anisotropic hyperelastic material model, the variable **cmname** can be tested for different material names inside user subroutine **VUANISOHYPER\_INV**, as illustrated below:

```
if (cmname(1:4) .eq. 'MAT1') then
    call VUANISOHYPER_INV1(argument_list)
else if (cmname(1:4) .eq. 'MAT2') then
    call VUANISOHYPER_INV2(argument_list)
end if
```

**VUANISOHYPER\_INV1** and **VUANISOHYPER\_INV2** are the actual subroutines containing the anisotropic hyperelastic models for each material **MAT1** and **MAT2**, respectively. Subroutine **VUANISOHYPER\_INV** merely acts as a directory here. The argument list can be the same as that used in subroutine **VUANISOHYPER\_INV**. The material names must be in uppercase characters since **cmname** is passed in as an uppercase character string.

---

### **Example: Anisotropic hyperelastic model of Kaliske and Schmidt**

As an example of the coding of subroutine **VUANISOHYPER\_INV**, consider the model proposed by Kaliske and Schmidt (2005) for nonlinear anisotropic elasticity with two families of fibers. The strain energy function is given by a polynomial series expansion in the form

$$\begin{aligned}
U = & \frac{1}{D}(J - 1)^2 + \sum_{i=1}^3 a_i(\bar{I}_1 - 3)^i + \sum_{j=1}^3 b_j(\bar{I}_2 - 3)^j + \sum_{k=2}^6 c_k(\bar{I}_{4(11)} - 1)^k + \sum_{l=2}^6 d_l(\bar{I}_{5(11)} - 1)^l \\
& + \sum_{m=2}^6 e_m(\bar{I}_{4(22)} - 1)^m + \sum_{n=2}^6 f_n(\bar{I}_{5(22)} - 1)^n + \sum_{p=2}^6 g_p(\zeta_{12}\bar{I}_{4(12)} - \zeta_{12}^2)^p.
\end{aligned}$$

The code in subroutine **VUANISOHYPER\_INV** must return the derivatives of the strain energy function with respect to the scalar invariants, which are readily computed from the above expression. In this example auxiliary functions are used to facilitate enumeration of pseudo-invarinats of type  $\bar{I}_{4(\alpha\beta)}$  and  $\bar{I}_{5(\alpha\beta)}$ , as well as for indexing into the array of second derivatives using symmetric storage. The subroutine would be coded as follows:

```

subroutine vuanisohyper_inv (
C Read only -
*      nblock, nFiber, nInv,
*      jElem, kIntPt, kLayer, kSecPt,
*      cmname,
*      nstatev, nfieldv, nprops,
*      props, tempOld, tempNew, fieldOld, fieldNew,
*      stateOld, sInvariant, zeta,
C     Write only -
*      uDev, duDi, d2uDiDi,
*      stateNew )
C
include 'vaba_param.inc'
C
dimension props(nprops),
*      tempOld(nblock),
*      fieldOld(nblock,nfieldv),
*      stateOld(nblock,nstatev),
*      tempNew(nblock),
*      fieldNew(nblock,nfieldv),
*      sInvariant(nblock,nInv),
*      zeta(nblock,nFiber*(nFiber-1)/2),
*      uDev(nblock), duDi(nblock,*),
*      d2uDiDi(nblock,*),
*      stateNew(nblock,nstatev)
C
character*80 cmname
C
parameter ( zero = 0.d0, one = 1.d0, two = 2.d0,
*      three = 3.d0, four = 4.d0, five = 5.d0, six = 6.d0 )

```

```
C
C      Kaliske energy function (3D)
C
C      Read material properties
d=props(1)
dinv = one / d
a1=props(2)
a2=props(3)
a3=props(4)
b1=props(5)
b2=props(6)
b3=props(7)
c2=props(8)
c3=props(9)
c4=props(10)
c5=props(11)
c6=props(12)
d2=props(13)
d3=props(14)
d4=props(15)
d5=props(16)
d6=props(17)
e2=props(18)
e3=props(19)
e4=props(20)
e5=props(21)
e6=props(22)
f2=props(23)
f3=props(24)
f4=props(25)
f5=props(26)
f6=props(27)
g2=props(28)
g3=props(29)
g4=props(30)
g5=props(31)
g6=props(32)
C
      do k = 1, nblock
         Udev(k) = zero
C      Compute Udev and 1st and 2nd derivatives w.r.t invariants
C      - I1
```

```

        bi1 = sInvariant(k,1)
        term = bi1-three
        Udev(k) = Udev(k)
        *      + a1*term + a2*term**2 + a3*term**3
        duDi(k,1) = a1 + two*a2*term + three*a3*term**2
        d2uDiDi(k,indx(1,1)) = two*a2 + three*two*a3*term
C      - I2
        bi2 = sInvariant(k,2)
        term = bi2-three
        Udev(k) = Udev(k)
        *      + b1*term + b2*term**2 + b3*term**3
        duDi(k,2) = b1 + two*b2*term + three*b3*term**2
        d2uDiDi(k,indx(2,2)) = two*b2 + three*two*b3*term
C      - I3 (=J)
        bi3 = sInvariant(k,3)
        term = bi3-one
        duDi(k,3) = two*dinv*term
        d2uDiDi(k,indx(3,3)) = two*dinv
C      - I4(11)
        nI411 = indxInv4(1,1)
        bi411 = sInvariant(k,nI411)
        term = bi411-one
        Udev(k) = Udev(k)
        *      + c2*term**2 + c3*term**3 + c4*term**4
        *      + c5*term**5 + c6*term**6
        duDi(k,nI411) =
        *      two*c2*term
        *      + three*c3*term**2
        *      + four*c4*term**3
        *      + five*c5*term**4
        *      + six*c6*term**5
        d2uDiDi(k,indx(nI411,nI411)) =
        *      two*c2
        *      + three*two*c3*term
        *      + four*three*c4*term**2
        *      + five*four*c5*term**3
        *      + six*five*c6*term**4
C      - I5(11)
        nI511 = indxInv5(1,1)
        bi511 = sInvariant(k,nI511)
        term = bi511-one
        Udev(k) = Udev(k)

```

```

*
*      + d2*term**2 + d3*term**3 + d4*term**4
*
*      + d5*term**5 + d6*term**6
duDi(k,nI511) =
*
*          two*d2*term
*
*          + three*d3*term**2
*
*          + four*d4*term**3
*
*          + five*d5*term**4
*
*          + six*d6*term**5
d2uDiDi(k,indx(nI511,nI511)) =
*
*          two*d2
*
*          + three*two*d3*term
*
*          + four*three*d4*term**2
*
*          + five*four*d5*term**3
*
*          + six*five*d6*term**4
C   - I4(22)
    nI422 = indxInv4(2,2)
    bi422 = sInvariant(k,nI422)
    term = bi422-one
    Udev(k) = Udev(k)
*
*          + e2*term**2 + e3*term**3 + e4*term**4
*
*          + e5*term**5 + e6*term**6
duDi(k,nI422) =
*
*          two*e2*term
*
*          + three*e3*term**2
*
*          + four*e4*term**3
*
*          + five*e5*term**4
*
*          + six*e6*term**5
d2uDiDi(k,indx(nI422,nI422)) =
*
*          two*e2
*
*          + three*two*e3*term
*
*          + four*three*e4*term**2
*
*          + five*four*e5*term**3
*
*          + six*five*e6*term**4
C   - I5(22)
    nI522 = indxInv5(2,2)
    bi522 = sInvariant(k,nI522)
    term = bi522-one
    Udev(k) = Udev(k)
*
*          + f2*term**2 + f3*term**3 + f4*term**4
*
*          + f5*term**5 + f6*term**6
duDi(k,nI522) =
*
*          two*f2*term

```

```

*
*      + three*f3*term**2
*      + four*f4*term**3
*      + five*f5*term**4
*      + six*f6*term**5
d2uDIDi(k,indx(nI522,nI522)) =
*
*      two*f2
*      + three*two*f3*term
*      + four*three*f4*term**2
*      + five*four*f5*term**3
*      + six*five*f6*term**4
C      - I4(12)
nI412 = indxInv4(1,2)
bi412 = sInvariant(k,nI412)
term = zeta(k,1)*(bi412-zeta(k,1))
Udev(k) = Udev(k)
*
*      + g2*term**2 + g3*term**3
*      + g4*term**4 + g5*term**5
*      + g6*term**6
duDi(k,nI412) = zeta(k,1) * (
*
*      two*g2*term
*      + three*g3*term**2
*      + four*g4*term**3
*      + five*g5*term**4
*      + six*g6*term**5 )
d2uDIDi(k,indx(nI412,nI412)) = zeta(k,1)**2 * (
*
*      two*g2
*      + three*two*g3*term
*      + four*three*g4*term**2
*      + five*four*g5*term**3
*      + six*five*g6*term**4 )

C      end do
C
C      return
C
C      Function to map index from Square to Triangular storage
C      of symmetric matrix
C
integer function indx( i, j )
include 'vaba_param.inc'
ii = min(i,j)

```

```
jj = max(i,j)
indx = ii + jj*(jj-1)/2
return
end

C
C Function to generate enumeration of scalar
C Pseudo-Invariants of type 4

C integer function indxInv4( i, j )
include 'vaba_param.inc'
ii = min(i,j)
jj = max(i,j)
indxInv4 = 4 + jj*(jj-1) + 2*(ii-1)
return
end

C
C Function to generate enumeration of scalar
C Pseudo-Invariants of type 5
C

integer function indxInv5( i, j )
include 'vaba_param.inc'
ii = min(i,j)
jj = max(i,j)
indxInv5 = 5 + jj*(jj-1) + 2*(ii-1)
return
end
```

---

## Additional reference

- Kaliske, M., and J. Schmidt, “Formulation of Finite Nonlinear Anisotropic Elasticity,” CADFEM GmbH Infoplanner 2/2005, vol. 2, pp. 22–23, 2005.

## 1.2.9 VUANISOHYPER\_STRAIN: User subroutine to define anisotropic hyperelastic material behavior based on Green strain.

**Product:** Abaqus/Explicit

### References

---

- “Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual
- \*ANISOTROPIC HYPERELASTIC
- “**VUANISOHYPER\_INV** and **VUANISOHYPER\_INV**,” Section 4.1.13 of the Abaqus Verification Manual

### Overview

---

User subroutine **VUANISOHYPER\_STRAIN**:

- can be used to define the strain energy potential of anisotropic hyperelastic materials as a function of the components of the Green strain tensor;
- will be called for blocks of material calculation points for which the material definition contains user-defined anisotropic hyperelastic behavior with Green strain-based formulation (“Anisotropic hyperelastic behavior,” Section 19.5.3 of the Abaqus Analysis User’s Manual);
- can use and update solution-dependent state variables;
- can use any field variables that are passed in; and
- requires that the values of the derivatives of the strain energy density function be defined with respect to the components of the modified Green strain tensor and the volume ratio.

### Component ordering in tensors

---

The component ordering depends upon whether the tensor is second or fourth order.

#### Symmetric second-order tensors

For symmetric second-order tensors, such as the modified Green strain tensor, there are **ndir+nshr** components; the component order is given as a natural permutation of the indices of the tensor. The direct components are first and then the indirect components, beginning with the 12-component. For example, a stress tensor contains **ndir** direct stress components and **nshr** shear stress components, which are passed in as

Component	2-D Case	3-D Case
1	$\bar{\varepsilon}_{11}^G$	$\bar{\varepsilon}_{11}^G$
2	$\bar{\varepsilon}_{22}^G$	$\bar{\varepsilon}_{22}^G$

Component	2-D Case	3-D Case
3	$\bar{\varepsilon}_{33}^G$	$\bar{\varepsilon}_{33}^G$
4	$\bar{\varepsilon}_{12}^G$	$\bar{\varepsilon}_{12}^G$
5		$\bar{\varepsilon}_{23}^G$
6		$\bar{\varepsilon}_{31}^G$

The shear strain components are stored as tensor components and not as engineering components.

### Symmetric fourth-order tensors

For symmetric fourth-order tensors, such as the deviatoric elasticity tensor  $\partial^2 U / \partial \bar{\varepsilon}_{ij}^G \partial \bar{\varepsilon}_{kl}^G$ , there are  $(\text{ndir} + \text{nshr}) * (\text{ndir} + \text{nshr} + 1) / 2$  independent components. These components are ordered using the following triangular storage scheme:

Component	2-D Case	3-D Case
1	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{11}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{11}^G$
2	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{22}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{22}^G$
3	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{22}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{22}^G$
4	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{33}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{33}^G$
5	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{33}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{33}^G$
6	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{33}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{33}^G$
7	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{12}^G$
8	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{12}^G$
9	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{12}^G$
10	$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{12}^G$	$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{12}^G$
11		$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{23}^G$
12		$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{23}^G$
13		$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{23}^G$
14		$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{23}^G$
15		$\partial^2 U / \partial \bar{\varepsilon}_{23}^G \partial \bar{\varepsilon}_{23}^G$
16		$\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{31}^G$

Component	2-D Case	3-D Case
17		$\partial^2 U / \partial \bar{\varepsilon}_{22}^G \partial \bar{\varepsilon}_{31}^G$
18		$\partial^2 U / \partial \bar{\varepsilon}_{33}^G \partial \bar{\varepsilon}_{31}^G$
19		$\partial^2 U / \partial \bar{\varepsilon}_{12}^G \partial \bar{\varepsilon}_{31}^G$
20		$\partial^2 U / \partial \bar{\varepsilon}_{23}^G \partial \bar{\varepsilon}_{31}^G$
21		$\partial^2 U / \partial \bar{\varepsilon}_{31}^G \partial \bar{\varepsilon}_{31}^G$

If  $Q$  denotes the component number of term  $\partial^2 U / \partial \bar{\varepsilon}_{ij}^G \partial \bar{\varepsilon}_{kl}^G$  in the above table and  $M$  and  $N$  (with  $M \leq N$ ) denote the component numbers of  $\bar{\varepsilon}_{ij}^G$  and  $\bar{\varepsilon}_{kl}^G$ , respectively, in the table for second-order tensors,  $Q$  is given by the relationship  $Q = M + N \times (N - 1)/2$ . For example, consider the term  $\partial^2 U / \partial \bar{\varepsilon}_{11}^G \partial \bar{\varepsilon}_{23}^G$ . The component numbers for  $\bar{\varepsilon}_{11}^G$  and  $\bar{\varepsilon}_{23}^G$  are  $M = 1$  and  $N = 5$ , respectively, giving  $Q = 1 + (5 \times 4)/2 = 11$ .

## Special consideration for shell elements

---

When **VUANISOHYPER\_STRAIN** is used to define the material response of shell elements, Abaqus/Explicit cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element's transverse shear stiffness. See "Shell section behavior," Section 26.6.4 of the Abaqus Analysis User's Manual, for guidelines on choosing this stiffness.

## Material point deletion

---

Material points that satisfy a user-defined failure criterion can be deleted from the model (see "User-defined mechanical material behavior," Section 23.8.1 of the Abaqus Analysis User's Manual). You must specify the state variable number controlling the element deletion flag when you allocate space for the solution-dependent state variables, as explained in "User-defined mechanical material behavior," Section 23.8.1 of the Abaqus Analysis User's Manual. The deletion state variable should be set to a value of one or zero in **VUANISOHYPER\_STRAIN**. A value of one indicates that the material point is active, and a value of zero indicates that Abaqus/Explicit should delete the material point from the model by setting the stresses to zero. The structure of the block of material points passed to user subroutine **VUANISOHYPER\_STRAIN** remains unchanged during the analysis; deleted material points are not removed from the block. Abaqus/Explicit will "freeze" the values of the strains passed to **VUANISOHYPER\_STRAIN** for all deleted material points; that is, the strain values remain constant after deletion is triggered. Once a material point has been flagged as deleted, it cannot be reactivated.

## User subroutine interface

---

```

subroutine vuanisohyper_strain(
C Read only (unmodifiable) variables -
  1 nblock,jElem,kIntPt,kLayer,kSecPt,cmname,

```

```

2 ndir,nshr,nstatev,nfieldv,nprops,
3 props,tempOld,tempNew,fieldOld,fieldNew,
4 stateOld,ebar,detu,
C Write only (modifiable) variables -
4 udev,duDe,duDj,
5 d2uDeDe,d2uDjDj,d2uDeDj,
6 stateNew)
C
      include 'vaba_param.inc'
C
      dimension jElem(nblock),
1  props(nprops),
2  tempOld(nblock),
3  fieldOld(nblock,nfieldv),
4  stateOld(nblock,nstatev),
5  tempNew(nblock),
6  fieldNew(nblock,nfieldv),
7  ebar(nblock,ndir+nshr), detu(nblock),
8  uDev(nblock),
9  duDe(nblock,ndir+nshr), duDj(nblock),
* d2uDeDe(nblock,(ndir+nshr)*(ndir+nshr+1)/2),
1  d2uDjDj(nblock),
2  d2uDeDj(nblock,ndir+nshr),
3  stateNew(nblock,nstatev)
C
      character*80 cmname
C

      do 100 km = 1,nblock
      user coding
100 continue

      return
      end

```

---

**Variables to be defined****udev (nblock)**

$\hat{U}_{dev}$ , the deviatoric part of the strain energy density of the primary material response. This quantity is needed only if the current material definition also includes Mullins effect (see “Mullins effect,” Section 19.6.1 of the Abaqus Analysis User’s Manual).

**duDe (nblock,ndir+nshr)**

Derivatives of strain energy potential with respect to the components of the modified Green strain tensor,  $\partial U / \partial \bar{\varepsilon}_{ij}^G$ .

**duDj (nblock,ndir+nshr)**

Derivatives of strain energy potential with respect to volume ratio,  $\partial U / \partial J$ .

**d2uDeDe (nblock,(ndir+nshr)\*(ndir+nshr+1)/2)**

Second derivatives of strain energy potential with respect to the components of the modified Green strain tensor (using triangular storage),  $\partial^2 U / \partial \bar{\varepsilon}_{ij}^G \partial \bar{\varepsilon}_{kl}^G$ .

**d2uDjDj (nblock)**

Second derivatives of strain energy potential with respect to volume ratio,  $\partial^2 U / \partial J^2$ .

**d2uDeDj (nblock,ndir+nshr)**

Cross derivatives of strain energy potential with respect to components of the modified Green strain tensor and volume ratio,  $\partial^2 U / \partial \bar{\varepsilon}_{ij}^G \partial J^2$ .

**stateNew (nblock,nstatev)**

State variables at each material point at the end of the increment. You define the size of this array by allocating space for it (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information).

---

## Variables passed in for information

**nblock**

Number of material points to be processed in this call to **VUANISOHYPER\_STRAIN**.

**jElem (nblock)**

Array of element numbers.

**kIntPt**

Integration point number.

**kLayer**

Layer number (for composite shells).

**kSecPt**

Section point number within the current layer.

**cmname**

User-specified material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, you should not use “ABQ\_” as the leading string for **cmname**.

## VUANISOHYPER\_STRAIN

### **ndir**

Number of direct components in a symmetric tensor.

### **nshr**

Number of indirect components in a symmetric tensor.

### **nstatev**

Number of user-defined state variables that are associated with this material type (you define this as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

### **nfieldv**

Number of user-defined external field variables.

### **nprops**

User-specified number of user-defined material properties.

### **props (nprops)**

User-supplied material properties.

### **tempOld (nblock)**

Temperatures at each material point at the beginning of the increment.

### **tempNew (nblock)**

Temperatures at each material point at the end of the increment.

### **fieldOld (nblock, nfieldv)**

Values of the user-defined field variables at each material point at the beginning of the increment.

### **fieldNew (nblock, nfieldv)**

Values of the user-defined field variables at each material point at the end of the increment.

### **stateOld (nblock, nstatev)**

State variables at each material point at the beginning of the increment.

### **ebar (nblock, ndir+nshr)**

Modified Green strain tensor,  $\bar{\epsilon}^G$ , at each material point at the end of the increment.

### **detu (nblock)**

$J$ , determinant of deformation gradient (volume ratio) at the end of the increment.

---

### **Example: Using more than one user-defined anisotropic hyperelastic material model**

To use more than one user-defined anisotropic hyperelastic material model, the variable **cmmname** can be tested for different material names inside user subroutine **VUANISOHYPER\_STRAIN**, as illustrated below:

```

if (cmname(1:4) .eq. 'MAT1') then
    call VUANISOHYPER_STRAIN1(argument_list)
else if (cmname(1:4) .eq. 'MAT2') then
    call VUANISOHYPER_STRAIN2(argument_list)
end if

```

**VUANISOHYPER\_STRAIN1** and **VUANISOHYPER\_STRAIN2** are the actual subroutines containing the anisotropic hyperelastic models for each material **MAT1** and **MAT2**, respectively. Subroutine **VUANISOHYPER\_STRAIN** merely acts as a directory here. The argument list can be the same as that used in subroutine **VUANISOHYPER\_STRAIN**. The material names must be in uppercase characters since **cmname** is passed in as an uppercase character string.

#### Example: Orthotropic Saint-Venant Kirchhoff model

---

As a simple example of the coding of subroutine **VUANISOHYPER\_STRAIN**, consider the generalization to anisotropic hyperelasticity of the Saint-Venant Kirchhoff model. The strain energy function of the Saint-Venant Kirchhoff model can be expressed as a quadratic function of the Green strain tensor,  $\varepsilon^G$ , as

$$U(\varepsilon^G) = \frac{1}{2} \varepsilon^G : \mathbf{D} : \varepsilon^G,$$

where  $\mathbf{D}$  is the fourth-order elasticity tensor. The derivatives of the strain energy function with respect to the Green strain are given as

$$\frac{\partial U}{\partial \varepsilon^G} = \mathbf{D} : \varepsilon^G,$$

$$\frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G} = \mathbf{D}.$$

However, subroutine **VUANISOHYPER\_STRAIN** must return the derivatives of the strain energy function with respect to the modified Green strain tensor,  $\bar{\varepsilon}^G$ , and the volume ratio,  $J$ , which can be accomplished easily using the following relationship between  $\varepsilon^G$ ,  $\bar{\varepsilon}^G$ , and  $J$ :

$$\varepsilon^G = J^{\frac{2}{3}} \bar{\varepsilon}^G + \frac{1}{2} (J^{\frac{2}{3}} - 1) \mathbf{I},$$

where  $\mathbf{I}$  is the second-order identity tensor. Thus, using the chain rule we find

$$\frac{\partial U}{\partial \bar{\varepsilon}^G} = J^{\frac{2}{3}} \frac{\partial U}{\partial \varepsilon^G},$$

$$\frac{\partial U}{\partial J} = \frac{\partial \varepsilon^G}{\partial J} : \frac{\partial U}{\partial \varepsilon^G},$$

## VUANISOHYPER\_STRAIN

$$\frac{\partial^2 U}{\partial \bar{\varepsilon}^G \partial \bar{\varepsilon}^G} = J^{\frac{4}{3}} \frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G},$$

$$\frac{\partial^2 U}{\partial J^2} = \frac{\partial^2 \varepsilon^G}{\partial J^2} : \frac{\partial U}{\partial \varepsilon^G} + \frac{\partial \varepsilon^G}{\partial J} : \frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G} : \frac{\partial \varepsilon^G}{\partial J},$$

$$\frac{\partial^2 U}{\partial \bar{\varepsilon}^G \partial J} = \frac{2}{3J} J^{\frac{2}{3}} \frac{\partial U}{\partial \varepsilon^G} + J^{\frac{2}{3}} \frac{\partial^2 U}{\partial \varepsilon^G \partial \varepsilon^G} : \frac{\partial \varepsilon^G}{\partial J},$$

where

$$\frac{\partial \varepsilon^G}{\partial J} = \frac{2}{3J} J^{\frac{2}{3}} (\bar{\varepsilon}^G + \frac{1}{2} \mathbf{I})$$

and

$$\frac{\partial^2 \varepsilon^G}{\partial J^2} = -\frac{1}{3J} \frac{\partial \varepsilon^G}{\partial J}.$$

In this example an auxiliary function is used to facilitate indexing into a fourth-order symmetric tensor. The subroutine would be coded as follows:

```

subroutine vuanisohyper_strain (
C Read only -
*      nblock,
*      jElem, kIntPt, kLayer, kSecPt,
*      cmname,
*      ndir, nshr, nstatev, nfieldv, nprops,
*      props, tempOld, tempNew, fieldOld, fieldNew,
*      stateOld, ebar, detu,
C Write only -
*      uDev, duDe, duDj,
*      d2uDeDe, d2uDjDj, d2uDeDj,
*      stateNew )
C
     include 'vaba_param.inc'
C
     dimension props(nprops),
*      tempOld(nblock),
*      fieldOld(nblock,nfieldv),
*      stateOld(nblock,nstatev),
*      tempNew(nblock),
*      fieldNew(nblock,nfieldv),
*      ebar(nblock,ndir+nshr), detu(nblock),

```

```

*  uDev(nblock), duDe(nblock,ndir+nshr), duDj(nblock),
*  d2uDeDe(nblock,*), d2uDjDj(nblock),
*  d2uDeDj(nblock,ndir+nshr),
*  stateNew(nblock,nstatev)

C
      character*80 cmname
C
      parameter( half = 0.5d0, one = 1.d0, two = 2.d0,
*            third = 1.d0/3.d0, twothds = 2.d0/3.d0, four = 4.d0,
*            dinv = 0.d0 )
C
C      Orthotropic Saint-Venant Kirchhoff strain energy function
C      (3D)
C
      D1111 = props(1)
      D1122 = props(2)
      D2222 = props(3)
      D1133 = props(4)
      D2233 = props(5)
      D3333 = props(6)
      D1212 = props(7)
      D1313 = props(8)
      D2323 = props(9)
C
      do k = 1, nblock
C
      d2UdE11dE11 = D1111
      d2UdE11dE22 = D1122
      d2UdE11dE33 = D1133
      d2UdE22dE11 = d2UdE11dE22
      d2UdE22dE22 = D2222
      d2UdE22dE33 = D2233
      d2UdE33dE11 = d2UdE11dE33
      d2UdE33dE22 = d2UdE22dE33
      d2UdE33dE33 = D3333
      d2UdE12dE12 = D1212
      d2UdE13dE13 = D1313
      d2UdE23dE23 = D2323
C
      xpow = exp ( log(detu(k)) * twothds )
      detuInv = one / detu(k)
C

```

## VUANISOHYPER\_STRAIN

```

E11 = xpow * ebar(k,1) + half * ( xpow - one )
E22 = xpow * ebar(k,2) + half * ( xpow - one )
E33 = xpow * ebar(k,3) + half * ( xpow - one )
E12 = xpow * ebar(k,4)
E23 = xpow * ebar(k,5)
E13 = xpow * ebar(k,6)

C
term1 = twothds * xpow * detuInv
dE11Dj = term1 * ( E11 + half )
dE22Dj = term1 * ( E22 + half )
dE33Dj = term1 * ( E33 + half )
dE12Dj = term1 * E12
dE13Dj = term1 * E13
dE23Dj = term1 * E23
term2 = - third * detuInv
d2E11DjDj = term2 * dE11Dj
d2E22DjDj = term2 * dE22Dj
d2E33DjDj = term2 * dE33Dj
d2E12DjDj = term2 * dE12Dj
d2E13DjDj = term2 * dE13Dj
d2E23DjDj = term2 * dE23Dj

C
dUdE11 = d2UdE11dE11 * E11
*           + d2UdE11dE22 * E22
*           + d2UdE11dE33 * E33
dUdE22 = d2UdE22dE11 * E11
*           + d2UdE22dE22 * E22
*           + d2UdE22dE33 * E33
dUdE33 = d2UdE33dE11 * E11
*           + d2UdE33dE22 * E22
*           + d2UdE33dE33 * E33
dUdE12 = two * d2UdE12dE12 * E12
dUdE13 = two * d2UdE13dE13 * E13
dUdE23 = two * d2UdE23dE23 * E23

C
U = half * ( E11*dUdE11 + E22*dUdE22 + E33*dUdE33 )
*           + E12*dUdE12 + E13*dUdE13 + E23*dUdE23
uDev(k) = U

C
duDe(k,1) = xpow * dUdE11
duDe(k,2) = xpow * dUdE22
duDe(k,3) = xpow * dUdE33

```

```

duDe(k, 4) = xpow * dUdE12
duDe(k, 5) = xpow * dUdE23
duDe(k, 6) = xpow * dUdE13
C
xpow2 = xpow * xpow
C Only update nonzero components
d2uDeDe(k, indx(1,1)) = xpow2 * d2UdE11dE11
d2uDeDe(k, indx(1,2)) = xpow2 * d2UdE11dE22
d2uDeDe(k, indx(2,2)) = xpow2 * d2UdE22dE22
d2uDeDe(k, indx(1,3)) = xpow2 * d2UdE11dE33
d2uDeDe(k, indx(2,3)) = xpow2 * d2UdE22dE33
d2uDeDe(k, indx(3,3)) = xpow2 * d2UdE33dE33
d2uDeDe(k, indx(4,4)) = xpow2 * d2UdE12dE12
d2uDeDe(k, indx(5,5)) = xpow2 * d2UdE23dE23
d2uDeDe(k, indx(6,6)) = xpow2 * d2UdE13dE13
*
duDj(k) = dUdE11*dE11Dj + dUdE22*dE22Dj + dUdE22*dE22Dj
*      + two * ( dUdE12*dE12Dj + dUdE13*dE13Dj
*                  + dUdE23*dE23Dj )
d2uDjDj(k) = dUdE11*d2E11DjDj+dUdE22*d2E22DjDj
*      +dUdE22*d2E22DjDj
*      + two*(dUdE12*d2E12DjDj+dUdE13*d2E13DjDj
*                  +dUdE23*d2E23DjDj)
*      + d2UdE11dE11 * dE11Dj * dE11Dj
*      + d2UdE22dE22 * dE22Dj * dE22Dj
*      + d2UdE33dE33 * dE33Dj * dE33Dj
*      + two * ( d2UdE11dE22 * dE11Dj * dE22Dj
*                  + d2UdE11dE33 * dE11Dj * dE33Dj
*                  + d2UdE22dE33 * dE22Dj * dE33Dj )
*      + four * ( d2UdE12dE12 * dE12Dj * dE12Dj
*                  d2UdE13dE13 * dE13Dj * dE13Dj
*                  d2UdE23dE23 * dE23Dj * dE23Dj )
*
d2uDeDj(k,1) = term1 * dUdE11 + xpow * (
*          d2UdE11dE11 * dE11Dj
*          + d2UdE11dE22 * dE22Dj
*          + d2UdE11dE33 * dE33Dj )
d2uDeDj(k,2) = term1 * dUdE22 + xpow * (
*          d2UdE22dE11 * dE11Dj
*          + d2UdE22dE22 * dE22Dj
*          + d2UdE22dE33 * dE33Dj )
d2uDeDj(k,3) = term1 * dUdE33 + xpow * (
*          + d2UdE33dE11 * dE11Dj

```

## VUANISOHYPER\_STRAIN

```
*           + d2UdE33dE22 * dE22Dj
*           + d2UdE33dE33 * dE33Dj )
d2uDeDj(k,4) = term1 * dUdE12
*           + xpow * two * d2UdE12dE12 * dE12Dj
d2uDeDj(k,5) = term1 * dUdE23
*           + xpow * two * d2UdE23dE23 * dE23Dj
d2uDeDj(k,6) = term1 * dUdE13
*           + xpow * two * d2UdE13dE13 * dE13Dj
end do
C
      return
end
C
integer function indx( i, j )
C
include 'vaba_param.inc'
C
C Function to map index from Square to Triangular storage
C of symmetric matrix
C
ii = min(i,j)
jj = max(i,j)
C
indx = ii + jj*(jj-1)/2
C
return
end
```

### 1.2.10 VUEL: User subroutine to define an element.

**Product:** Abaqus/Explicit

*WARNING: This feature is intended for advanced users only. Its use in all but the simplest test examples will require considerable coding by the user/developer. “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual, should be read before proceeding.*

#### References

---

- “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual
- “User-defined element library,” Section 29.16.2 of the Abaqus Analysis User’s Manual
- “UEL,” Section 1.1.24
- \*UEL PROPERTY
- \*USER ELEMENT

#### Overview

---

User subroutine **VUEL**:

- will be called for each element that is of a general user-defined element type each time element calculations are required; and
- (or subroutines called by user subroutine **VUEL**) must perform all of the calculations for the element, appropriate to the current activity in the analysis.

#### User subroutine interface

---

```

SUBROUTINE VUEL(nblock,rhs,amass,dtimeStable,svars,nsvars,
1                 energy,
2                 nnod,ndofel,props,nprops,jprops,njprops,
3                 coords,mcrd,u,du,v,a,
4                 jtype,jElem,
5                 time,period,dtimeCur,dtimePrev,kstep,kinc,
6                 lflags,
7                 dMassScaleFactor,
8                 predef,npref,
9                 jdltyp, adlmg)
C
C           include 'vaba_param.inc'
C
C           operational code keys

```

```

        parameter ( jMassCalc          = 1,
*                  jIntForceAndDtStable = 2,
*                  jExternForce        = 3)

C      flag indices
parameter (iProcedure = 1,
*                  iNlgeom    = 2,
*                  iOpCode    = 3,
*                  nFlags     = 3)

C      energy array indices
parameter ( iElPd = 1,
*                  iElCd = 2,
*                  iElIe = 3,
*                  iElTs = 4,
*                  iElDd = 5,
*                  iElBv = 6,
*                  iElDe = 7,
*                  iElHe = 8,
*                  iElKe = 9,
*                  iElTh = 10,
*                  iElDmd = 11,
*                  iElDc = 12,
*                  nElEnergy = 12)

C      predefined variables indices
parameter ( iPredValueNew = 1,
*                  iPredValueOld = 2,
*                  nPred         = 2)

C      time indices
parameter (iStepTime   = 1,
*                  iTotalTime = 2,
*                  nTime       = 2)

dimension rhs(nblock,ndofel),amass(nblock,ndofel,ndofel),
1                  dtimestable(nblock),
2                  svars(nblock,nsvars),energy(nblock,nElEnergy),
3                  props(nprops),jprops(njprops),
4                  jElem(nblock),time(nTime),lflags(nFlags),
5                  coords(nblock,nnode,mcrd),
6                  u(nblock,ndofel), du(nblock,ndofel),

```

```

7      v(nblock,ndofel), a(nblock, ndofel),
8      dMassScaleFactor(nblock),
9      predef(nblock,nnode,npredef,nPred),
*      adlmag(nblock)

do kblock = 1,nblock
  user coding to define rhs, amass, dtimestable, svars and energy
end do

RETURN
END

```

## Variables to be defined

---

Some of the following arrays depend on the value of the **lflags** array.

### **rhs**

An array containing the contributions of each element to the right-hand-side vector of the overall system of equations. Depending on the settings of the **lflags** array, it contains either the internal force from the element or the external load calculated from the specified distributed loads.

### **amass**

An array containing the contribution of each element to the mass matrix of the overall system of equations.

All nonzero entries in **amass** should be defined. Moreover, the mass matrix must be symmetric. There are several other requirements that apply depending on the active degrees of freedom specified. These requirements are explained in detail below.

### **dtimestable**

A scalar value defining, for each element, the upper limit of the time increment for stability considerations. This would be the maximum time increment to be used in the subsequent increment for this element to be stable (to satisfy the Courant condition). This value depends strongly on the element formulation, and it is important that is computed appropriately.

### **svars**

An array containing the values of the solution-dependent state variables associated with each element. The number of such variables is **nsvars** (see below). You define the meaning of these variables.

This array is passed into **VUEL** containing the values of these variables at the start of the current increment. In most cases they should be updated to be the values at the end of the increment. In rare cases such an update is not required.

### **energy**

The array **energy** contains the values of the energy quantities associated with each element. The values in this array when **VUEL** is called are the element energy quantities at the start of the current

increment. They should be updated to the correct values at the end of the current increment; otherwise, plots of the energy balance for the entire model will not be accurate. Depending on the element formulation, many of these energies could be zero at all times. The entries in the array are as follows:

<b>energy (nblock, iElPd )</b>	Plastic dissipation.
<b>energy (nblock, iElCd)</b>	Creep dissipation.
<b>energy (nblock, iElIe)</b>	Internal energy.
<b>energy (nblock, iElTs)</b>	Transverse shear energy.
<b>energy (nblock, iElDd)</b>	Material damping dissipation.
<b>energy (nblock, iElBv)</b>	Bulk viscosity dissipation.
<b>energy (nblock, iElDe)</b>	Drill energy.
<b>energy (nblock, iElHe)</b>	Hourglass energy.
<b>energy (nblock, iElKe)</b>	Kinetic energy.
<b>energy (nblock, iElTh)</b>	Heat energy.
<b>energy (nblock, iElDmd)</b>	Damage dissipation.
<b>energy (nblock, iElDc )</b>	Distortion control energy.

## Variables passed in for information

---

### Arrays:

#### **props**

A floating point array containing the **nprops** real property values defined for use with each element processed. **nprops** is the user-specified number of real property values. See “Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual.

#### **jprops**

An integer array containing the **njprops** integer property values defined for use with each element processed. **njprops** is the user-specified number of integer property values. See “Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual.

#### **coords**

An array containing the original coordinates of the nodes of the element. **coords (kblock, k1 ,k2)** is the **k2**th coordinate of the **k1**th node of the **kblock** element.

**u, du, v, a**

Arrays containing the basic solution variables (displacements, rotations, temperatures, pressures, depending on the degree of freedom) at the nodes of the element. Values are provided as follows, illustrated below for the **k1**th degree of freedom of the **kblock** element:

<b>u (kblock, k1)</b>	Total value of the variables (such as displacements or rotations) at the end of the current increment.
<b>du (kblock, k1)</b>	Incremental values of the variables in the current increment.
<b>v (kblock, k1)</b>	Time rate of change of the variables (velocities, rates of rotation) at the midpoint of the increment.
<b>a (kblock, k1)</b>	Accelerations of the variables at the end of the current increment.

**jElem**

**jElem (kblock)** contains the element number for the **kblock** element.

**adlmag**

**adlmag (kblock)** is the total load magnitude of the load type **jdltyp** (integer identifying the load number for distributed load type *Un*) distributed load at the end of the current increment for distributed loads of type *Un*.

**predef**

An array containing the values of predefined field variables, such as temperature in an uncoupled stress/displacement analysis, at the nodes of the element (“Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual).

The second index, **k2**, indicates the local node number on the **kblock** element. The third index, **k3**, indicates the variable: the temperature is stored if the index is 1, and the predefined field variables are stored if the indices are greater than or equal to 2. The fourth index of the array, **k4**, is either 1 or 2, with 1 indicating the value of the field variable at the end of the increment and 2 indicating the value of the field variable at the beginning of the increment.

<b>predef (kblock, k2, 1, k4)</b>	Temperature.
<b>predef (kblock, k2, 2, k4)</b>	First predefined field variable.
<b>predef (kblock, k2, 3, k4)</b>	Second predefined field variable.
Etc.	Any other predefined field variable.

<b>predef(kblock,k2,k3,k4)</b>	Value of the ( <b>k3-1</b> ) th predefined field variable at the <b>k2</b> th node of the element at the beginning or the end of the increment.
<b>predef(kblock,k2,k3,1)</b>	Values of the variables at the end of the current increment.
<b>predef(kblock,k2,k3,2)</b>	Values of the variables at the beginning of the current increment.

**lflags**

An array containing the flags that define the current solution procedure and requirements for element calculations.

<b>lflags(iProcedure)</b>	Defines the procedure type. See “Results file output format,” Section 5.1.2 of the Abaqus Analysis User’s Manual, for the key used for each procedure.
<b>lflags(iNlgeom)=0</b>	Small-displacement analysis.
<b>lflags(iNlgeom)=1</b>	Large-displacement analysis (nonlinear geometric effects included in the step; see “General and linear perturbation procedures,” Section 6.1.2 of the Abaqus Analysis User’s Manual).
<b>lflags(iOpCode)=jMassCalc</b>	Define the mass matrix <b>amass</b> in the beginning of the analysis.
<b>lflags(iOpCode)=jIntForceAndDtStable</b>	Define the element internal force. Define the stable time increment as well.
<b>lflags(iOpCode)=jExternForce</b>	Define the distributed load effect on the external force associated with the element.

**dMassScaleFactor**

An array containing the mass scale factors for each element.

**time(iStepTime)**

Current value of step time.

**time (iTTotalTime)**  
 Current value of total time.

### Scalar parameters:

**nblock**  
 Number of user elements to be processed in this call to **VUEL**.

**dtimeCur**  
 Current time increment.

**dtimePrev**  
 Previous time increment.

**period**  
 Time period of the current step.

**ndofel**  
 Number of degrees of freedom in the elements processed.

**nsvars**  
 User-defined number of solution-dependent state variables associated with the element (“Defining the number of solution-dependent variables that must be stored within the element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**nprops**  
 User-defined number of real property values associated with the elements processed (“Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**njprops**  
 User-defined number of integer property values associated with the elements processed (“Defining the element properties” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**mcrd**  
**mcrd** is defined as the maximum of the user-defined maximum number of coordinates needed at any node point (“Defining the maximum number of coordinates needed at any nodal point” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual) and the value of the largest active degree of freedom of the user element that is less than or equal to 3. For example, if you specify that the maximum number of coordinates is 1 and the active degrees of freedom of the user element are 2, 3, and 6 **mcrd** will be 3. If you specify that the maximum number of coordinates is 2 and the active degree of freedom of the user element is 11, **mcrd** will be 2.

**nnode**

User-defined number of nodes on the elements (“Defining the number of nodes associated with the element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**jtype**

Integer defining the element type. This is the user-defined integer value  $n$  in element type  $VUn$  (“Assigning an element type key to a user-defined element” in “User-defined elements,” Section 29.16.1 of the Abaqus Analysis User’s Manual).

**kstep**

Current step number.

**kinc**

Current increment number.

**npreddef**

Number of predefined field variables, including temperature. For user elements Abaqus/Explicit uses one value for each field variable per node.

---

**VUEL conventions**

The solution variables (displacement, velocity, etc.) are arranged on a node/degree of freedom basis. The degrees of freedom of the first node are first, followed by the degrees of freedom of the second node, etc. The degrees of freedom that will be updated automatically in Abaqus/Explicit are: 1–3 (displacements), 4–6 (rotations), 8 (pressure), and 11 (temperature). Depending on the procedure type (see below), only some of the degrees of freedom listed above will be updated. Other degrees of freedom will not be updated by the time integration procedure in Abaqus/Explicit and, hence, should not be used.

The mass matrix defined in user subroutine **VUEL** must be symmetric. In addition, the following requirements apply:

- The mass matrix entries associated with the translational degrees of freedom for a particular node must be diagonal. Moreover, these diagonal entries must be equal to each other.
- There must be no coupling (off-diagonal) entries specified between degrees of freedom of different kinds. For example, you cannot specify nonzero mass matrix entries to couple the translational degrees of freedom to the rotational degrees of freedom.
- There must be no coupling (off-diagonal) entries specified between degrees of freedom belonging to different nodes.

You must be using appropriate lumping techniques to provide a mass matrix that follows these requirements. For the rotational degrees of freedom at a particular node in three-dimensional analyses, you can specify a fully populated symmetric  $3 \times 3$  inertia tensor.

---

**Usage with general nonlinear procedures**

The following illustrates the use in explicit dynamic procedures:

**Direct-integration explicit dynamic analysis (`lflags (iProcedure)=17`)**

- Automatic updates for degrees of freedom 1–6, 8, and 11.
- The governing equations are as described in “Explicit dynamic analysis,” Section 6.3.3 of the Abaqus Analysis User’s Manual.
- Coding for the operational code `lflags (iOpCode)=jExternForce` is optional.

**Transient fully coupled thermal-stress analysis (`lflags (iProcedure)=74`)**

- Automatic updates for degrees of freedom 1–6 and 8.
- The governing equations are as described in “Fully coupled thermal-stress analysis in Abaqus/Explicit” in “Fully coupled thermal-stress analysis,” Section 6.5.4 of the Abaqus Analysis User’s Manual.
- Coding for the operational code `lflags (iOpCode)=jExternForce` is optional.

**Example: Structural user element**

---

A structural user element has been created to demonstrate the usage of subroutine **VUEL**. These user-defined elements are applied in a number of analyses. The following excerpt is from the verification problem that invokes the structural user element in an explicit dynamic procedure:

```
*USER ELEMENT, NODES=2, TYPE=VU1, PROPERTIES=4, COORDINATES=3,
VARIABLES=12
1, 2, 3
*ELEMENT, TYPE=VU1
101, 101, 102
*ELGEN, ELSET=VUTRUSS
101, 5
*UEL PROPERTY, ELSET=VUTRUSS
0.002, 2.1E11, 0.3, 7200.
```

The user element consists of two nodes that are assumed to lie parallel to the *x*-axis. The element behaves similarly to a linear truss element. The supplied element properties are the cross-sectional area, Young’s modulus, Poisson’s ratio, and density, respectively.

The next excerpt shows the listing of the subroutine. The user subroutine has been coded for use in an explicit dynamic analysis. The names of the verification input files associated with the subroutine and these procedures can be found in “**VUEL**,” Section 4.1.32 of the Abaqus Verification Manual.

```
subroutine vuel(
*      nblock,
*      rhs, amass, dtimeStable,
*      svars, nsvars,
*      energy,
*      nnodes, ndofel,
```

```

*      props,nprops,
*      jprops,njprops,
*      coords,ncrd,
*      u,du,v,a,
*      jtype,jElem,
*      time,period,dtimeCur,dtimePrev,kstep,kinc,lfflags,
*      dMassScaleFactor,
*      predef,npref,
*      ndload,adlmag)

include 'vaba_param.inc'

c     operation code
parameter ( jMassCalc          = 1,
*              jIntForceAndDtStable = 4)

c     flags
parameter ( iProcedure = 1,
*                  iNlgeom    = 2,
*                  iOpCode    = 3,
*                  nFlags     = 3)

c     procedure flags
parameter ( jDynExplicit = 17 )

c     time
parameter ( iStepTime   = 1,
*                  iTotTime    = 2,
*                  nTime       = 2)

c     energies
parameter ( iElPd = 1,
*                  iElCd = 2,
*                  iElIe = 3,
*                  iElTs = 4,
*                  iElDd = 5,
*                  iElBv = 6,
*                  iElDe = 7,
*                  iElHe = 8,
*                  iElKe = 9,
*                  iElTh = 10,
*                  iElDmd = 11,

```

```

*           iElDc = 12,
*           nElEnergy = 12)

parameter (factorStable = 0.99d0)
parameter ( zero = 0.d0, half = 0.5d0, one = 1.d0, two=2.d0 )
c
dimension rhs(nblock,ndofel), amass(nblock,ndofel,ndofel),
*      dtimeStable(nblock),
*      svars(nblock,nsvars), energy(nblock,nElEnergy),
*      props(nprops), jprops(njprops),
*      jElem(nblock), time(nTime), l(nFlags),
*      coords(nblock,nnode,ncrd), u(nblock,ndofel),
*      du(nblock,ndofel), v(nblock,ndofel), a(nblock, ndofel),
*      predef(nblock, nnode, npredef, nPred), adlmag(nblock),
*      dMassScaleFactor(nblock)

c      Notes:
c      Define only nonzero entries; the arrays to be defined have
c      been zeroed out just before this call

if (jtype .eq. 1001 .and.
*     lflags(iProcedure) .eq. jDynExplicit) then

area0 = props(1)
eMod  = props(2)
anu   = props(3)
rho   = props(4)

eDampTra    = zero
amassFact0  = half*area0*rho

if ( lflags(iOpCode) .eq. jMassCalc ) then
do kblock = 1, nblock
      use original distance to compute mass
      alenX0 = (coords(kblock,2,1) - coords(kblock,1,1))
      alenY0 = (coords(kblock,2,2) - coords(kblock,1,2))
      alenZ0 = (coords(kblock,2,3) - coords(kblock,1,3))
      alen0 = sqrt(alenX0*alenX0 + alenY0*alenY0 +
*                  alenZ0*alenZ0)
      am0   = amassFact0*alen0
      amass(kblock,1,1) = am0

```

```

        amass(kblock,2,2) = am0
        amass(kblock,3,3) = am0
        amass(kblock,4,4) = am0
        amass(kblock,5,5) = am0
        amass(kblock,6,6) = am0

        end do
    else if ( lflags(iOpCode) .eq.
    *           jIntForceAndDtStable) then
        do kblock = 1, nblock
            alenX0 = (coords(kblock,2,1) - coords(kblock,1,1))
            alenY0 = (coords(kblock,2,2) - coords(kblock,1,2))
            alenZ0 = (coords(kblock,2,3) - coords(kblock,1,3))
            alen0 = sqrt(alenX0*alenX0 + alenY0*alenY0 +
    *                   alenZ0*alenZ0)
            vol0 = area0*alen0
            amElem0 = two*amassFact0*alen0

            alenX = alenX0
            *                   + (u(kblock,4) - u(kblock,1))
            alenY = alenY0
            *                   + (u(kblock,5) - u(kblock,2))
            alenZ = alenZ0
            *                   + (u(kblock,6) - u(kblock,3))
            alen = sqrt(alenX*alenX + alenY*alenY + alenZ*alenZ)
            area   = vol0/alen
            ak     = area*eMod/alen

c         stable time increment for translations
c         dtimStable(kblock) = factorStable*sqrt(amElem0/ak)

c         force = E * logarithmic strain *current area
c         strainLog = log(alen/alen0)
c         fElastra = eMod*strainLog*area

        forceTra = fElastra

c         assemble internal load in RHS
        rhs(kblock,1) = -forceTra
        rhs(kblock,4) = forceTra

c         internal energy calculation

```

```
    alenOld = svars(kblock,1)
    fElasTraOld = svars(kblock,2)

    energy(kblock, iElle) = energy(kblock, iElle) +
    half*(fElasTra+fElasTraOld)*(alen - alenOld)

c      update state variables
    svars(kblock,1) = alen
    svars(kblock,2) = fElasTra

    end do
    end if

    end if
c
    return
end
```



## 1.2.11 VUFIELD: User subroutine to specify predefined field variables.

**Product:** Abaqus/Explicit

### References

---

- “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual
- \*FIELD

### Overview

---

User subroutine **VUFIELD**:

- allows you to prescribe predefined field variables at the nodes of a model—the predefined field variables at a node can be updated individually, or a number of field variables at the nodes can be updated simultaneously;
- can be called for blocks of nodes for which the field variable values are defined in the subroutine;
- ignores any field variable values specified directly;
- can be used to modify field variable values read from a results file; and
- can be used in conjunction with user subroutine **VUSDFLD** such that the field variables that are passed in from **VUFIELD** and interpolated to the material points can be modified (such changes are local to material point values, and nodal field variable values remain unaffected).

### Updating field variables

---

Two different methods are provided for updating field variables.

#### Individual variable updates

By default, only one field variable is updated at a time for given nodes or a given node set in user subroutine **VUFIELD**. The user subroutine is called whenever a current value of a field variable is needed for the nodes that are listed in the field variable definition. This method is ideal for cases in which the field variables are independent of each other.

#### Simultaneous variable updates

User subroutine **VUFIELD** can also be used to update multiple field variables simultaneously for given nodes or a given node set. This method is well-suited for cases in which there are dependencies between some of the field variables. In this case you must specify the number of field variables to be updated simultaneously, and the user subroutine will be called each time the field variable values are needed.

---

User subroutine interface

---

```
SUBROUTINE VUFIELD(FIELD, NBLOCK, NFIELD, KFIELD, NCOMP,
1                      KSTEP, KINC, JNODEID, TIME,
2                      COORDS, U, V, A)

C
INCLUDE 'VABA_PARAM.INC'

C      indices for the time array TIME
PARAMETER( i_ufld_Current    = 1,
*              i_ufld_Increment = 2,
*              i_ufld_Period     = 3,
*              i_ufld_Total       = 4 )

C      indices for the coordinate array COORDS
PARAMETER( i_ufld_CoordX = 1,
*                  i_ufld_CoordY = 2,
*                  i_ufld_CoordZ = 3 )

C      indices for the displacement array U
PARAMETER( i_ufld_SpaDisplX = 1,
*                  i_ufld_SpaDisplY = 2,
*                  i_ufld_SpaDisplZ = 3,
*                  i_ufld_RotDisplX = 4,
*                  i_ufld_RotDisplY = 5,
*                  i_ufld_RotDisplZ = 6,
*                  i_ufld_AcoPress   = 7,
*                  i_ufld_Temp        = 8 )

C      indices for the velocity array V
PARAMETER( i_ufld_SpaVelX   = 1,
*                  i_ufld_SpaVelY   = 2,
*                  i_ufld_SpaVelZ   = 3,
*                  i_ufld_RotVelX   = 4,
*                  i_ufld_RotVelY   = 5,
*                  i_ufld_RotVelZ   = 6,
*                  i_ufld_DAcoPress = 7,
*                  i_ufld_DTemp      = 8 )

C      indices for the acceleration array A
PARAMETER( i_ufld_SpaAccelX = 1,
```

```

*      i_ufld_SpaAccelY = 2,
*      i_ufld_SpaAccelZ = 3,
*      i_ufld_RotAccelX = 4,
*      i_ufld_RotAccelY = 5,
*      i_ufld_RotAccelZ = 6,
*      i_ufld_DDAcoPress = 7,
*      i_ufld_DDTemp     = 8 )
C
DIMENSION FIELD (NBLOCK,NCOMP,NFIELD)
DIMENSION JNODEID (NBLOCK), TIME (4), COORDS (3,NBLOCK)
DIMENSION U (8,NBLOCK), V (8,NBLOCK), A (8,NBLOCK)
C
user coding to define FIELD

RETURN
END

```

## Variable to be defined

---

### **FIELD (NBLOCK , NCOMP , NFIELD)**

Array of field variable values at a collective number of nodes **NBLOCK** (see **NBLOCK** below). When updating one field variable at a time, only the value of the specified field variable **KFIELD** must be returned. In this case **NFIELD** is passed into user subroutine **VUFIELD** with a value of 1, and **FIELD** is thus dimensioned as **FIELD (NBLOCK , NCOMP , 1)**. When updating all field variables simultaneously, the values of the specified number of field variables must be returned. In this case **FIELD** is dimensioned as **FIELD (NBLOCK , NCOMP , NFIELD)**, where **NFIELD** is the number of field variables specified and **KFIELD**, which is set to -1, has no meaning.

If fields are applied to nodes that are not part of pipe, beam, or shell elements, only one value of each field variable is required (**NCOMP**=1). Otherwise, the number of values to be returned depends on the mode of temperature and field variable input selected for the beam or shell section. It should also be noted that **VUFIELD** does not allow the use of values at the reference surface with gradients for either shell or beam elements. Temperature or field values at the section points through the thickness are assumed to be constant. However, you can employ gradients by specifying explicitly temperature or field values at the section points.

Because field variables can also be defined directly, it is important to understand the hierarchy used in situations with conflicting information (see “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual).

When the array **FIELD** is passed into user subroutine **VUFIELD**, it will contain either the field variable values from the previous increment or those values obtained from the results file if this method was used. You can then modify these values within this subroutine.

---

**Variables passed in for information****NBLOCK**

User-specified number of nodes to be processed in this call to **VUFIELD**. The value is equal to the total number of nodes given in a node set when the optional parameter BLOCKING on the \*FIELD option is omitted or is set to NO. When the parameter BLOCKING is set to YES, **NBLOCK** is equal to a predefined number set in Abaqus/Explicit. You can also modify **NBLOCK** by setting any desirable value through BLOCKING= $n$ .

**NFIELD**

User-specified number of field variables to be updated. The default value is 1.

**KFIELD**

User-specified field variable number. This variable is meaningful only when updating individual field variables at a time; otherwise, the value is set to -1.

**NCOMP**

Maximum number of section values required for any node in the model. When fields are applied to nodes that are part of pipe, beam, or shell elements, **VUFIELD** is invoked in two passes: the first pass with **NCOMP** passed in with the value of 1, and the second pass with **NCOMP** passed in with the value equal to the total number of section points minus 1.

**KSTEP**

Current step number.

**KINC**

Increment number for step **KSTEP**.

**JNODEUID (NBLOCK)**

Array for user-defined node numbers. This array is dimensioned based on the size of **NBLOCK**, and the contained node numbers are identical to those defined in the input file. You can perform additional interdependent field variable operations by using nodal indices stored in this array.

**TIME (4)**

Array for information of analysis time. You can retrieve any time information from this array by using the parameters given above. **TIME(i\_ufld\_Current)** stores the current analysis time, **TIME(i\_ufld\_Increment)** gives the time increment at this instance, **TIME(i\_ufld\_Period)** is the time period of the current step, and **TIME(i\_ufld\_Total)** is the total analysis time up to this point. You can use this time information to perform possible time-dependent field variable operations.

**COORDS (3 , NBLOCK)**

Coordinates for nodes in the array **JNODEUID**. This array stores current coordinates of nodes in which the order of coordinates stored corresponds to the order of nodes listed in the array **JNODEUID**. The

coordinates can be retrieved by using the parameters given above. You can make use of **COORDS** to define possible position-dependent field variable operations.

**U(8,NBLOCK) , V(8,NBLOCK) , and A(8,NBLOCK)**

Arrays containing solution variables of displacements, rotations, temperatures, and pressures and their corresponding temporal derivatives. The order in which these solutions are stored follows the order defined in the array **JNODEUID**. For a specific node its solution variables can be retrieved by using the parameter indices given above. Depending on the degrees of freedom, some solution variables are not valid for a given node.



## 1.2.12 VUFLUIDEXCH: User subroutine to define the mass flow rate/heat energy flow rate for fluid exchange.

**Product:** Abaqus/Explicit

### References

---

- “Fluid exchange definition,” Section 11.6.3 of the Abaqus Analysis User’s Manual
- \*FLUID EXCHANGE
- \*FLUID EXCHANGE ACTIVATION
- \*FLUID EXCHANGE PROPERTY

### Overview

---

User subroutine **VUFLUIDEXCH**:

- can be used to define mass flow rate and/or heat energy flow rate for fluid exchange;
- can be used when built-in fluid exchange property types cannot satisfactorily model the mass/heat energy flow;
- can use and update solution-dependent state variables;
- can use any field variables that are passed in; and
- requires that the derivatives of mass/heat energy flow rates be defined with respect to pressure and temperature in the primary and secondary fluid cavities.

### Conventions for defining mass flow/heat energy flow rate

---

A positive mass/heat energy flow rate indicates flow from the primary fluid cavity to the secondary fluid cavity. A negative value for mass flow rate will be ignored if the fluid exchange is between a cavity and its environment.

### User subroutine interface

---

```

subroutine vufluidexch(
C Read only (unmodifiable)variables -
  1      nstatev, nfieldv, nprops,
  2      stepTime, totalTime, dt,
  3      jCavType, fluExchName, effArea, amplitude,
  4      props, lExchEnv, pcavNew, pcavOld,
  5      ctempNew, ctempOld, cvol, cmass,
  6      rMix, CpMix, DCpDtemp,
  7      field, stateOld,
```

```

C Write only (modifiable) variables
    8      stateNew, rMassRate, rEneRate,
    9      DMassRateDPcav, DMassRateDTemp,
    *      DEneRateDPcav, DEneRateDTemp)
c
c      include 'vaba_param.inc'
c
c      dimension props(nprops),
1      pcavNew(2), pcavOld(2),
2      ctempNew(2), ctempOld(2), cvol(2), cmass(2),
3      rMix(2), CpMix(2), dCpDtemp(2),
4      field(nfieldv),
5      stateOld(nstatev), stateNew(nstatev),
6      DMassRateDPcav(2), DMassRateDTemp(2),
7      DEneRateDPcav(2), DEneRateDTemp(2)

c      Fluid cavity type
parameter( iHydraulic      = 1,
*                  iAdiabaticGas   = 2,
*                  iIsothermalGas = 3)

character*80 fluExchName

c      User coding to calculate mass flow rate,
c      heat energy flow rate and its derivatives with respect
c      to fluid cavity pressure and temperature.

return
end

```

## Variables to be defined

---

### **rMassRate**

Mass flow rate. The mass flow rate is negative if the flow is into the primary cavity.

### **DMassRateDPcav (2)**

Derivative of mass flow rate with respect to pressure in primary and secondary fluid cavities.

### **DMassRateDTemp (2)**

Derivative of mass flow rate with respect to temperature in primary and secondary fluid cavities.

### **rEneRate**

Heat energy flow rate. The energy flow rate is negative if the flow is into the primary cavity.

**DEneRateDPcav (2)**

Derivative of heat energy flow rate with respect to pressure in primary and secondary fluid cavities.

**DEneRateDTemp (2)**

Derivative of heat energy flow rate with respect to temperature in primary and secondary fluid cavities.

**Variable that can be updated**

---

**stateNew (nstatev)**

State variable for fluid exchange at the end of the increment. You define the size of this array by allocating space for it (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information).

**Variables passed in for information**

---

**nstatev**

Number of user-defined state variables that are associated with this fluid exchange (you define this as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined fluid exchange properties required to define mass/heat energy flow rate.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**dt**

Time increment size.

**jCavType**

Indicator of fluid cavity type: 1 for fluid cavity with hydraulic fluids, 2 for fluid cavity with adiabatic gases, and 3 for fluid cavity with isothermal gases.

**fluExchName**

User-specified fluid exchange name.

**effArea**

Effective area for fluid exchange.

**amplitude**

Current value of the amplitude referenced for this fluid exchange. You must multiply the flow rates by the current amplitude value within the user subroutine if the amplitude is required.

**props (nprop)**

User-defined fluid exchange properties.

**lExchEnv**

The fluid exchange is to the environment if **lExchEnv**=1 and to another fluid cavity if **lExchEnv**=0.

**pcavNew (2)**

Pressure in primary and secondary fluid cavities at the end of the increment.

**pcavOld (2)**

Pressure in primary and secondary fluid cavities at the beginning of the increment.

**ctempNew (2)**

Temperature in primary and secondary fluid cavities at the end of the increment.

**ctempOld (2)**

Temperature in primary and secondary fluid cavities at the beginning of the increment.

**cvol (2)**

Volume of primary and secondary fluid cavities.

**cmass (2)**

Mass of fluid in primary and secondary fluid cavities.

**rMix (2)**

Gas constant of mixture in primary and secondary fluid cavities.

**CpMix (2)**

Specific heat of mixture in primary and secondary fluid cavities.

**DCpDttemp (2)**

Derivative of specific heat with respect to temperature for primary and secondary fluid cavities.

**field(nfieldv)**

Field variables at orifice.

**stateOld (nstatev)**

State variables for fluid exchange at the beginning of the increment.

## 1.2.13 VUFLUIDEXCHEFFAREA: User subroutine to define the effective area for fluid exchange.

**Product:** Abaqus/Explicit

### References

---

- “Fluid exchange definition,” Section 11.6.3 of the Abaqus Analysis User’s Manual
- \*FLUID EXCHANGE

### Overview

---

User subroutine **VUFLUIDEXCHEFFAREA**:

- can be used to define an effective area for fluid exchange that depends on the material state in the underlying elements on the fluid exchange surface;
- will be called for blocks of material calculation points on the fluid exchange surface;
- can be used only if the specified surface over which the fluid exchange occurs is a surface defined over membrane elements; and
- can be used with any fluid exchange property type.

### Defining effective area

---

The contribution of each material point can be defined as a function of:

- the original area associated with the material point;
- the current material state in the underlying elements; and
- the temperature and pressure in the primary fluid cavity and the secondary fluid cavity or environment.

The effective area for fabric materials can depend on the nominal strain in the yarn directions and the change in angle between the two yarn directions, as well as the current angle between the two yarn directions. For nonfabric materials the effective area can depend on the material point strain.

### User subroutine interface

---

```

subroutine vufluidexcheffarea(
C Read only (unmodifiable) variables -
  1      nblock, nprop, props,
  2      stepTime, totalTime, fluExchName,
  3      cMatName, lFabric, braidAngle,
  4      strain, origArea,
```

```

      5      pcav, ctemp,
C Write only (modifiable) variables
      6      effArea)

c
      include 'vaba_param.inc'
c
      parameter (ndir = 3, nshr=1)
c
c      pointers for retrieving fabric constitutive strains
      parameter( iFiberStrain1    = 1,
*              iFiberStrain2    = 2,
*              iFiberChangeAng  = 4)
c
      dimension props(nprop),
1      braidAngle(nblock),
2      strain(nblock, ndir+nshr),
3      origArea(nblock),
4      pcav(2),ctemp(2),
5      effArea(nblock)

      character*80 fluExchName, cMatName

c      do k = 1, nblock
c          User coding to update effArea(k) = area associated with
c          material point contributing to area for fluid exchange
c          (leakage).
c      end do

      return
end

```

---

**Variable to be defined****effArea(nblock)**

Area associated with the material point contributing to the total effective area for fluid exchange. The subroutine is called with **effArea** set to the current area associated with the material point and should be updated to reflect the area that contributes to fluid exchange.

---

**Variables passed in for information****nBlock**

Number of material points to be processed in this call to **VUFLUIDEXCHEFFAREA**.

**nprop**

User-specified number of user-defined fluid exchange properties required to define the effective area.

**props (nprop)**

User-defined fluid exchange properties.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**fluExchName**

User-specified fluid exchange name.

**cMatName**

User-specified material name associated with material points processed in this call.

**lFabric**

Flag indicating whether the subroutine is called for material points on the fluid exchange surface with a fabric material (**lFabric**=1 if fabric material, **lFabric**=0 otherwise).

**braidAngle (nblock)**

Angle in radians between the two yarn directions for fabric materials.

**strain (nblock, ndir+nshr)**

Fabric constitutive strains (nominal strain in the yarn directions and change in angle between the two yarn directions) or strains for nonfabric materials at current location.

**origArea (nblock)**

Original area associated with current material point.

**pcav (2)**

Absolute pressure in primary and secondary (or ambient) fluid cavities at the start of the increment.

**ctemp (2)**

Temperature in primary and secondary (or ambient) fluid cavities at the start of the increment.



**1.2.14 VUHARD: User subroutine to define the yield surface size and hardening parameters for isotropic plasticity or combined hardening models.**

**Product:** Abaqus/Explicit

## References

---

- “Classical metal plasticity,” Section 20.2.1 of the Abaqus Analysis User’s Manual
- “Models for metals subjected to cyclic loading,” Section 20.2.2 of the Abaqus Analysis User’s Manual
- \*CYCLIC HARDENING
- \*PLASTIC
- “Deformation of a sandwich plate under CONWEP blast loading,” Section 9.1.8 of the Abaqus Example Problems Manual
- “**VUHARD**,” Section 4.1.34 of the Abaqus Verification Manual

## Overview

---

User subroutine **VUHARD**:

- is called at all material points of elements for which the material definition includes user-defined isotropic hardening or cyclic hardening for metal plasticity;
- can be used to define a material’s isotropic yield behavior;
- can be used to define the size of the yield surface in a combined hardening model;
- can include material behavior dependent on field variables or state variables; and
- requires that the derivatives of the yield stress (or yield surface size in combined hardening models) be defined with respect to the appropriate independent variables, such as strain, strain rate, and temperature.

## User subroutine interface

---

```

        subroutine vuhard(
C Read only -
        *      nblock,
        *      jElem, kIntPt, kLayer, kSecPt,
        *      lAnneal, stepTime, totalTime, dt, cmname,
        *      nstatev, nfieldv, nprops,
        *      props, tempOld, tempNew, fieldOld, fieldNew,
        *      stateOld,
        *      eqps, eqpsRate,
C Write only -

```

```

*      yield, dyieldDtemp, dyieldDeqps,
*      stateNew )

C
    include 'vaba_param.inc'
C
    dimension props(nprops), tempOld(nblock), tempNew(nblock),
1    fieldOld(nblock,nfieldv), fieldNew(nblock,nfieldv),
2    stateOld(nblock,nstatev), eqps(nblock), eqpsRate(nblock),
3    yield(nblock), dyieldDtemp(nblock), dyieldDeqps(nblock,2),
4    stateNew(nblock,nstatev), jElem(nblock)
C
    character*80 cmname
C
    do 100 km = 1,nblock
        user coding
100 continue
C
    return
end

```

## Variables to be defined

---

### **yield(nblock)**

Array containing the yield stress (for isotropic plasticity) or yield surface size (for combined hardening) at the material points.

### **dyieldDeqps (nblock,1)**

Array containing the derivative of the yield stress or yield surface size with respect to the equivalent plastic strain at the material points.

### **dyieldDeqps (nblock,2)**

Array containing the derivative of the yield stress with respect to the equivalent plastic strain rate at the material points. This quantity is not used with the combined hardening model.

### **dyieldDtemp (nblock)**

Array containing the derivative of the yield stress or yield surface size with respect to temperature at the material points. This quantity is required only in adiabatic and fully coupled temperature-displacement analyses.

### **stateNew(nblock,nstatev)**

Array containing the state variables at the material points at the end of the increment. The allocation of this array is described in “Solution-dependent state variables” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

---

## Variables passed in for information

**nblock**

Number of material points to be processed in this call to **VUHARD**.

**jElem(nblock)**

Array of element numbers.

**kIntPt**

Integration point number.

**kLayer**

Layer number (for composite shells).

**kSecPt**

Section point number within the current layer.

**lanneal**

Flag indicating whether the routine is being called during an annealing process. **lanneal**=0 indicates that the routine is being called during a normal mechanics increment. **lanneal**=1 indicates that this is an annealing process and the internal state variables, **stateNew**, should be reinitialized if necessary. Abaqus/Explicit will automatically set the stresses, stretches, and state to a value of zero during the annealing process.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**dt**

Time increment size.

**cmname**

Material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, “ABQ\_” should not be used as the leading string for **cmname**.

**nstatev**

Number of user-defined state variables that are associated with this material type (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**tempOld(nblock)**

Temperatures at the material points at the beginning of the increment.

**tempNew(nblock)**

Temperatures at the material points at the end of the increment.

**fieldOld(nblock,nfieldv)**

Values of the user-defined field variables at the material points at the beginning of the increment.

**fieldNew(nblock,nfieldv)**

Values of the user-defined field variables at the material points at the end of the increment.

**stateOld(nblock,nstatev)**

State variables at the material points at the beginning of the increment.

**eqps(nblock)**

Equivalent plastic strain at the material points.

**eqpsRate(nblock)**

Equivalent plastic strain rate at the material points.

## 1.2.15 VUINTER: User subroutine to define the interaction between contact surfaces.

**Product:** Abaqus/Explicit

### References

---

- “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual
- \*SURFACE INTERACTION
- “**VUINTER**,” Section 4.1.35 of the Abaqus Verification Manual

### Overview

---

User subroutine **VUINTER**:

- can be used to define the mechanical and thermal interaction between contacting surfaces;
- must provide the entire definition of the interaction between the contacting surfaces;
- can use and update solution-dependent state variables; and
- must be used with the penalty contact constraint algorithm.

### Terminology

---

The use of user subroutine **VUINTER** requires familiarity with the following terminology.

### Surface node numbers

The “surface node number” refers to the position of a particular node in the list of nodes on the surface. For example, there are **nSlvNod** nodes on the slave surface. Number  $n, n = 1, 2, \dots, n_{SlvNod}$ , is the surface node number of the  $n$ th node in this list; **jSlvUid**( $n$ ) is the user-defined global number of this node. An Abaqus/Explicit model can be defined in terms of an assembly of part instances (see “Defining an assembly,” Section 2.9.1 of the Abaqus Analysis User’s Manual). In such models a node number in **jSlvUid** is an internally generated node number. If the original node number and part instance name are required, call the utility routine **VGETPARTINFO** (see “Obtaining part information,” Section 2.1.5).

### Local coordinate system

---

The array **alocaldir** defines the direction cosines of a local coordinate system for each slave node. The first local direction corresponds to the contact normal direction from the perspective of the slave node. For a two-dimensional **VUINTER** model the second local direction is the tangent direction defined by the cross product of the vector into the plane of the model (0., 0., -1.0) and the slave normal. For a three-dimensional **VUINTER** model the second and third local directions correspond to two orthogonal tangent directions  $t_1$  and  $t_2$ , which are set as follows:

- If the master surface is a cylindrical analytical surface, the second local direction corresponds to the generator direction (see “Analytical rigid surface definition,” Section 2.3.4 of the Abaqus Analysis User’s Manual), and the third local direction is the cross product of the first and second local directions.
- If the master surface is an analytical surface of revolution, the third local direction corresponds to the hoop direction, and the second local direction is the cross product of the third and first local directions.
- If the master surface is a three-dimensional, element-based surface, the tangent directions are based on the slave normal, using the standard convention for calculating surface tangents (see “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual).

For the two cases listed above involving three-dimensional analytical surfaces, the local tangent directions will reflect a rotation of the master surface. For the last case (three-dimensional, element-based master surface) the tangent directions may not follow the rotation of either the master or slave surfaces; for example, the local system would remain fixed with respect to the global system if a slave node and its surrounding facets rotate about an axes parallel to the slave normal.

The  $2 \times 2$  array stored in **drot** for each slave node represents the incremental rotation of the tangent directions within the tangent plane corresponding to the tracked point of a three-dimensional master surface. (This incremental rotation array is equal to a unit matrix if **nDir** is equal to 2.) This incremental rotation matrix is provided so that vector- or tensor-valued state variables defined within the tangent plane can be rotated in this subroutine. For example, the second and third components of the **rdisp** array (i.e., the relative slip components) are rotated by this amount before **VUINTER** is called. However, as already mentioned, the rotation of the tangent directions may not reflect a physical rotation of the master or slave surface.

### **Conventions for heat flux and stress**

---

A positive flux indicates heat flowing into a surface, and a negative flux denotes heat leaving the surface. Flux must be specified for both surfaces, and they need not be equal and opposite so that effects such as frictional dissipation and differential surface heating can be modeled.

A positive normal stress denotes a pressure directed into the surface (opposite the local normal direction). Positive shear stresses denote shear tractions in the direction of the local surface tangents.

### **User subroutine interface**

---

```

subroutine vuinter(
  C Write only
    1 sfd, scd, spd, svd,
  C Read/Write -
    2 stress, fluxSlv, fluxMst, sed, statev,
  C Read only -
    3 kStep, kInc, nFacNod, nSlvNod, nMstNod, nSurfDir,
    4 nDir, nStateVar, nProps, nTemp, nPred, numDefTfv,
```

```

5 jSlvUid, jMstUid, jConMstid, timStep, timGlb,
6 dTimCur, surfInt, surfSlv, surfMst,
7 rdisp, drdisp, drot, stiffDflt, condDflt,
8 shape, coordSlv, coordMst, alocaldir, props,
9 areaSlv, tempSlv, dtempSlv, preDefSlv, dpreDefSlv,
1 tempMst, dtempMst, preDefMst, dpreDefMst)
C
C           include `vaba_param.inc'
C
C           character*80 surfInt, surfSlv, surfMst
C
dimension props(nProps), statev(nStateVar,nSlvNod),
1 drot(2,2,nSlvNod), sed(nSlvNod), sfd(nSlvNod),
2 scd(nSlvNod), spd(nSlvNod), svd(nSlvNod),
3 rdisp(nDir,nSlvNod), drdisp(nDir,nSlvNod),
4 stress(nDir,nSlvNod), fluxSlv(nSlvNod),
5 fluxMst(nSlvNod), areaSlv(nSlvNod),
6 stiffDflt(nSlvNod), condDflt(nSlvNod),
7 alocaldir(nDir,nDir,nSlvNod), shape(nFacNod,nSlvNod),
8 coordSlv(nDir,nSlvNod), coordMst(nDir,nMstNod),
9 jSlvUid(nSlvNod), jMstUid(nMstNod),
1 jConMstid(nFacNod,nSlvNod), tempSlv(nSlvNod),
2 dtempSlv(nSlvNod), preDefSlv(nPred,nSlvNod),
3 dpreDefSlv(nPred,nSlvNod), tempMst(numDefTfv),
4 dtempMst(numDefTfv), preDefMst(nPred,numDefTfv),
5 dpreDefMst(nPred,numDefTfv)

user coding to define stress,
and, optionally, fluxSlv, fluxMst, statev, sed, sfd, scd, spd,
and svd

return
end

```

## Variable to be defined

---

### **stress(nDir, nSlvNod)**

On entry this array contains the stress at the interface during the previous time increment. It must be updated to the stress at the interface in the current time increment.

---

**Variables that can be updated****fluxSlv (nSlvNod)**

On entry this array contains the flux entering the slave surface during the previous time increment. It must be updated to the flux entering the slave surface during the current increment.

**fluxMst (nSlvNod)**

On entry this array contains the flux entering the master surface during the previous time increment. It must be updated to the flux entering the master surface during the current time increment.

**sfd (nSlvNod)**

This array can be updated to contain the increment in frictional dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLFD and have no effect on other solution variables.

**scd (nSlvNod)**

This array can be updated to contain the increment in creep dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLCD and have no effect on other solution variables.

**spd (nSlvNod)**

This array can be updated to contain the increment in plastic dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLPD and have no effect on other solution variables.

**svd (nSlvNod)**

This array can be updated to contain the increment in viscous dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLVD and have no effect on other solution variables.

**sed (nSlvNod)**

On entry this array contains the elastic energy density at the slave nodes at the beginning of the increment. It can be updated to contain the elastic energy density at the end of the current time increment. These values contribute to the output variable ALLSE and have no effect on other solution variables.

**statev (nstateVar, nSlvNod)**

This array contains the user-defined solution-dependent state variables for all the nodes on the slave surface. You define the size of this array (see “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual, for more information). This array will be passed in containing the values of these variables prior to the call to user subroutine **VUINTER**. If any of the solution-dependent state variables is being used in conjunction with the surface interaction, it must be updated in this subroutine.

---

## Variables passed in for information

**kStep**

Step number.

**kInc**

Increment number.

**nFacNod**

Number of nodes on each master surface facet. **nFacNod** is 2 for two-dimensional surfaces, and **nFacNod** is 4 for three-dimensional surfaces (the first and last nodes are the same for triangular facets). If the master surface is an analytical rigid surface, this variable is passed in as 0.

**nSlvNod**

Number of slave nodes.

**nMstNod**

Number of master surface nodes, if the master surface is made up of facets. If the master surface is an analytical rigid surface, this variable is passed in as 0.

**nSurfDir**

Number of tangent directions at the contact points (**nSurfDir** = **nDir** - 1).

**nDir**

Number of coordinate directions at the contact points. (In a three-dimensional model **nDir** will be 2 if the surfaces in the contact pair are two-dimensional analytical rigid surfaces or are formed by two-dimensional elements.)

**nStateVar**

Number of user-defined state variables.

**nProps**

User-specified number of property values associated with this surface interaction model.

**nTemp**

1 if the temperature is defined and 0 if the temperature is not defined.

**nPred**

Number of predefined field variables.

**numDefTfv**

Equal to **nSlvNod** if the master surface is made up of facets. If the master surface is an analytical rigid surface, this variable is passed in as 1.

**jSlvUid (nSlvNod)**

This array lists the user-defined global node numbers (or internal node numbers for models defined in terms of an assembly of part instances) of the nodes on the slave surface.

**jMstUid (nMstNod)**

This array lists the user-defined global node numbers (or internal node numbers for models defined in terms of an assembly of part instances) of the nodes on the master surface. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**jConMstid (nFacNod, nSlvNod)**

This array lists the surface node numbers of the master surface nodes that make up the facet onto which each slave node projects. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**timStep**

Value of step time.

**timGlb**

Value of total time.

**dtimCur**

Current increment in time from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ .

**surfInt**

User-specified surface interaction name, left justified.

**surfSlv**

Slave surface name.

**surfMst**

Master surface name.

**rdisp (nDir, nSlvNod)**

An array containing the relative positions between the two surfaces. The first component is the relative position of the slave node, with respect to the master surface, in the normal direction (a positive value indicates a penetration, and a negative value indicates a gap). The second and third components, if applicable, are the accumulated incremental relative tangential displacements of the slave node, measured from the beginning of the step in which the contact pair is defined. The local directions in which the relative displacements are defined are stored in **alocaldir**. If the master surface is an analytical surface, the elements in **rdisp** are set to **r\_MaxVal** for the slave nodes that are far from the master surface.

**drdisp (nDir, nSlvNod)**

An array containing the increments in relative positions between the two surfaces during the current time increment. If the master surface is an analytical surface, the elements in **drdisp** are set to **r\_MaxVal** for the slave nodes that are far from the master surface.

**drot (2, 2, nSlvNod)**

Rotation increment matrix. This matrix represents the incremental rotation of the local surface tangent directions for a three-dimensional surface. This rotation matrix for each slave node is defined as a unit matrix for two-dimensional surfaces. If the master surface is an analytical surface, the elements in **drot** are set to **r\_MaxVal** for the slave nodes that are far from the master surface.

**stiffDflt (nSlvNod)**

Values of the default penalty stiffnesses for each slave node (units of  $\text{FL}^{-3}$ ).

**condDflt (nSlvNod)**

Values of the default penalty conductances for each slave node (units of  $\text{J}\theta^{-1}\text{T}^{-1}$ ).

**shape (nFacNod, nSlvNod)**

For each contact point this array contains the shape functions of the nodes of its master surface facet, evaluated at the location of the contact point. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

**coordSlv (nDir, nSlvNod)**

Array containing the **nDir** components of the current coordinates of the slave nodes.

**coordMst (nDir, nMstNod)**

Array containing the **nDir** components of the current coordinates of the master nodes. If the master surface is an analytical rigid surface, this array is passed in as the coordinates of the contact points on the master surface.

**alocaldir (nDir, nDir, nSlvNod)**

Direction cosines of the local surface coordinate system. The first array index corresponds to the components of the local directions, and the second array index corresponds to the local direction number. The first direction (**alocaldir(1..nDir,1,...)**) is the normal to the surface. The second direction (**alocaldir(1..nDir,2,...)**) is the first surface tangent. For a three-dimensional surface, the third direction (**alocaldir(1..3,3,...)**) is the second surface tangent. If the master surface is an analytical rigid surface, the numbers in **alocaldir** are valid only if the corresponding parts in **rdisp** are valid (i.e., not equal to **r\_MaxVal**).

**props (nProps)**

User-specified vector of property values to define the behavior between the contacting surfaces.

**areaSlv (nSlvNod)**

Area associated with the slave nodes (equal to 1 for node-based surface nodes).

**tempSlv (nSlvNod)**

Current temperature at the slave nodes.

**dtempSlv (nSlvNod)**

Increment in temperature during the previous time increment at the slave nodes.

**preDefSlv (nPred, nSlvNod)**

Current user-specified predefined field variables at the slave nodes (initial values at the beginning of the analysis and current values during the analysis).

**dpreDefSlv (nPred, nSlvNod)**

Increment in the predefined field variables at the slave nodes during the previous time increment.

**tempMst (numDefTfv)**

Current temperature at the nearest points on the master surface.

**dtempMst (numDefTfv)**

Increment in temperature during the previous time increment at the nearest points on the master surface.

**preDefMst (nPred, numDefTfv)**

Current user-specified predefined field variables at the nearest points on the master surface (initial values at the beginning of the analysis and current values during the analysis).

**dpreDefMst (nPred, numDefTfv)**

Increment in the predefined field variables during the previous time increment at the nearest points on the master surface.

## 1.2.16 VUINTERACTION: User subroutine to define the contact interaction between surfaces with the general contact algorithm.

**Product:** Abaqus/Explicit

### References

---

- “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual
- \*SURFACE INTERACTION

### Overview

---

User subroutine **VUINTERACTION**:

- can be used to define the mechanical and thermal interaction between contact surfaces;
- must provide the entire definition of the interaction between the contact surfaces;
- can utilize a user-specified tracking thickness to determine the contact surfaces for node-to-surface contact;
- can use and update solution-dependent state variables for node-to-face contact; and
- must be used with the general contact algorithm.

### Local coordinate system

---

The array **dircos** defines the direction cosines of a local coordinate system for each slave node. The first local direction corresponds to the contact normal direction from the perspective of the slave node. The second and third local directions correspond to two orthogonal tangent directions  $t_1$  and  $t_2$ , which are set as follows:

- If the master surface is a cylindrical analytical surface, the second local direction corresponds to the generator direction (see “Analytical rigid surface definition,” Section 2.3.4 of the Abaqus Analysis User’s Manual), and the third local direction is the cross product of the first and second local directions.
- If the master surface is an analytical surface of revolution, the third local direction corresponds to the hoop direction, and the second local direction is the cross product of the third and first local directions.
- If the master surface is element-based, the tangent directions are based on the slave normal and the line connecting the first and third nodes on the master facet.

For the two cases listed above involving analytical surfaces, the local tangent directions will reflect a rotation of the master surface. For the last case (element-based master surface) the tangent directions follow the rotation of the master surface only approximately. The second tangent direction is constructed

such that it is perpendicular to the slave normal and the line going from the first to the third node on the master facet. The slave normal, the first tangent, and the second tangent form a right-handed system.

### Conventions for heat flux and stress

---

A positive flux indicates heat flowing into a surface, and a negative flux denotes heat leaving the surface. Flux must be specified for both surfaces, and they need not be equal and opposite so that effects such as frictional dissipation and differential surface heating can be modeled.

A positive normal stress denotes a pressure directed into the surface (opposite the local normal direction). Positive shear stresses denote shear tractions in the direction of the local surface tangents.

### User subroutine interface

---

```

        subroutine vuinteraction (
C Read/Write -
        *   stress, fluxSlv, fluxMst,
        *   state, sed,
C Write only -
        *   sfd, scd, spd, svd,
C Read only -
        *   nBlock, nBlockAnal, nBlockEdge,
        *   nNodState, nNodSlv, nNodMst, nDir,
        *   nStates, nProps, nTemp, nFields,
        *   jFlags, rData,
        *   surfInt, surfSlv, surfMst,
        *   jSlvUid, jMstUid, props,
        *   penetration, drDisp, dRot, dircos, stiffDef, conductDef,
        *   coordSlv, coordMst, areaSlv, shapeSlv, shapeMst,
        *   tempSlv, tempMst, dTempSlv, dTempMst,
        *   fieldSlv, fieldMst, dFieldSlv, dFieldMst )
C
        include `vaba_param.inc'
C
        dimension stress(nDir,nBlock),
        *   fluxSlv(nBlock),
        *   fluxMst(nBlock),
        *   state(nStates,nNodState,nBlock),
        *   sed(nBlock),
        *   sfd(nBlock),
        *   scd(nBlock),
        *   spd(nBlock),
        *   svd(nBlock),

```

```

*   jSlvUid(nNodSlv,nBlock),
*   jMstUid(nNodMst,nBlockAnal),
*   props(nProps),
*   penetration(nBlock),
*   drDisp(nDir,nBlock),
*   dRot(2,2,nBlock),
*   stiffDef(nBlock),
*   conductDef(nBlock),
*   dircos(nDir,nDir,nBlock),
*   coordSlv(nDir,nNodSlv,nBlock),
*   coordMst(nDir,nNodMst,nBlockAnal),
*   areaSlv(nBlock),
*   shapeSlv(nNodSlv,nBlockEdge),
*   shapeMst(nNodMst,nBlockAnal),
*   tempSlv(nBlock),
*   tempMst(nBlockAnal),
*   dTempSlv(nBlock),
*   dTempMst(nBlockAnal),
*   fieldSlv(nFields,nBlock),
*   fieldMst(nFields,nBlockAnal)
*   dFieldSlv(nFields,nBlock),
*   dFieldMst(nFields,nBlockAnal)

C
    parameter( iKStep      = 1,
*              iKIInc     = 2,
*              iLConType  = 3,
*              nFlags     = 3 )

C
    parameter( iTimStep    = 1,
*              iTimGlb     = 2,
*              iDTimCur   = 3,
*              iTrackThic = 4,
*              nData      = 4 )

C
    dimension jFlags(nFlags), rData(nData)

C
    character*80 surfInt, surfSlv, surfMst

C
user coding to define stress,
and, optionally, fluxSlv, fluxMst, state, sed, sfd, scd, spd,
and svd

C

```

```

return
end

```

## Variable to be defined

---

### **stress (nDir, nBlock)**

On entry this array contains the stress at the interface during the previous time increment. It must be updated to the stress at the interface in the current time increment.

## Variables that can be updated

---

### **fluxSlv (nBlock)**

On entry this array contains the flux entering the slave surface during the previous time increment. It must be updated to the flux entering the slave surface during the current increment.

### **fluxMst (nBlock)**

On entry this array contains the flux entering the master surface during the previous time increment. It must be updated to the flux entering the master surface during the current increment.

### **state (nStates, nNodState, nBlock)**

This array contains the user-defined solution-dependent state variables for all the nodes on the slave surface. The use of the state variables is applicable only for node-to-face contact. See “User-defined interfacial constitutive behavior,” Section 33.1.6 of the Abaqus Analysis User’s Manual, for more information on the size of this array. This array will be passed in containing the values of these variables prior to the call to user subroutine **VUINTERACTION**.

If any of the solution-dependent state variables is being used in conjunction with the interaction, it must be updated in this subroutine. These state variables need to be updated with care, as a slave node can be in contact with multiple master surfaces. Such a slave node may be passed into the user subroutine at a given increment multiple times, possibly on separate calls to the user subroutine, and you may end up advancing the state variables for that node multiple times for a single time increment. One trick to keep track of whether or not a node state is advanced is to use one of the state variables exclusively for this purpose. You could set that selected state variable to the current increment number and update the state only if it is not already set to the current increment number.

### **sed (nBlock)**

On entry this array contains the elastic energy density at the slave nodes at the beginning of the increment. It can be updated to contain the elastic energy density at the end of the current time increment. These values contribute to the output variable ALLSE and have no effect on other solution variables. The use of this variable is applicable only for node-to-face contact.

### **sfd (nBlock)**

This array can be updated to contain the increment in frictional dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLFD and have no effect on other solution variables. The use of this variable is applicable only for node-to-face contact.

**scd(nBlock)**

This array can be updated to contain the increment in creep dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLCD and have no effect on other solution variables. The use of this variable is applicable only for node-to-face contact.

**spd(nBlock)**

This array can be updated to contain the increment in plastic dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLPD and have no effect on other solution variables. The use of this variable is applicable only for node-to-face contact.

**svd(nBlock)**

This array can be updated to contain the increment in viscous dissipation at each node (units of energy per unit area). These values contribute to the output variables SFDR and ALLVD and have no effect on other solution variables. The use of this variable is applicable only for node-to-face contact.

---

**Variables passed in for information**

---

**nBlock**

Number of tracking points to be processed in this call to **VFRICITION**.

**nBlockAnal**

1 for analytical rigid master surface; **nBlock** otherwise.

**nBlockEdge**

**nBlock** for edge type slave surface; 1 otherwise.

**nNodState**

1 for node-to-face and node-to-analytical rigid surface contact; not applicable for edge-to-edge contact.

**nNodS1v**

1 for node-to-face and node-to-analytical rigid surface contact; 2 for edge-to-edge contact.

**nNodMst**

1 for analytical rigid master surface; 2 for edge-type master surface; 4 for facet-type master surface.

**nDir**

Number of coordinate directions at the tracking points (equal to 3).

**nStates**

Number of user-defined state variables.

**nProps**

User-specified number of property values associated with this friction model.

**nTemp**

1 if the temperature is defined and 0 if the temperature is not defined.

**nFields**

Number of predefined field variables.

**jFlag(1)**

Step number.

**jFlag(2)**

Increment number.

**jFlag(3)**

1 for node-to-face contact, 2 for edge-to-edge contact, and 3 for node-to-analytical rigid surface contact.

**rData(1)**

Value of step time.

**rData(2)**

Value of total time.

**rData(3)**

Current increment in time from  $t = t_{curr} - \Delta t$  to  $t = t_{curr}$ .

**rData(4)**

This variable contains the value of the tracking thickness specified for the surface interaction.

**surfInt**

User-specified surface interaction name, left justified.

**surfSlv**

Slave surface name, not applicable to general contact.

**surfMst**

Master surface name, not applicable to general contact.

**jSlvUid(nNodSlv,nBlock)**

This array lists the surface node numbers of the slave surface nodes that are tracked.

**jMstUid(nNodMst,nBlockAnal)**

This array lists the surface node numbers of the master surface nodes that make up the facet with which each slave node is tracked.

**props(nProps)**

User-specified vector of property values to define the interaction between the tracking surfaces.

**penetration(nBlock)**

The relative position of the slave node, with respect to the master surface, in the normal direction (a positive value indicates a penetration, and a negative value indicates a gap). If the master surface is an

analytical surface, the elements in **penetration** are set to **r\_MaxVal** for the slave nodes that are far from the master surface.

#### **drDisp (nDir, nBlock)**

An array containing the increments in relative positions between the two surfaces during the current time increment. If the master surface is an analytical surface, the elements in **drDisp** are set to **r\_MaxVal** for the slave nodes that are far from the master surface.

#### **dRot (2, 2, nBlock)**

This argument is currently undefined.

#### **stiffDef (nBlock)**

Values of the default penalty stiffnesses (stress per unit penetration) for each slave node (units of  $\text{FL}^{-3}$ ).

#### **conductDef (nBlock)**

Values of the default penalty conductances for each slave node (units of  $\text{J}\theta^{-1}\text{T}^{-1}$ ).

#### **dircos (nDir, nDir, nBlock)**

Direction cosines of the local surface coordinate system. The first array index corresponds to the components of the local directions, and the second array index corresponds to the local direction number. The first direction (**dircos (1..nDir, 1, ...)**) is the normal to the surface. The second direction (**dircos (1..nDir, 2, ...)**) is the first surface tangent. For a three-dimensional surface, the third direction (**dircos (1..3, 3, ...)**) is the second surface tangent. If the master surface is an analytical rigid surface, the numbers in **dircos** are valid only if the corresponding parts in **penetration** are valid (i.e., not equal to **r\_MaxVal**).

#### **coordSlv (nDir, nNodSlv, nBlock)**

Array containing the **nDir** components of the current coordinates of the slave nodes.

#### **coordMst (nDir, nNodMst, nBlockAnal)**

Array containing the **nDir** components of the current coordinates of the master nodes. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

#### **areaSlv (nBlock)**

Area associated with the slave nodes (equal to 1 for node-based surface nodes).

#### **shapeSlv (nNodSlv, nBlockEdge)**

For each contact point this array contains the shape functions of the nodes of its slave surface, evaluated at the location of the contact point.

#### **shapeMst (nNodMst, nBlockAnal)**

For each contact point this array contains the shape functions of the nodes of its master surface, evaluated at the location of the contact point. If the master surface is an analytical rigid surface, this array is passed in as a dummy array.

## VUINTERACTION

### **tempSlv(nBlock)**

Current temperature at the contact points on the slave surface.

### **tempMst(nBlockAnal)**

Current temperature at the contact points on the master surface.

### **dTempSlv(nBlock)**

Increment in the temperature during the previous time increment at the slave nodes.

### **dTempMst(nBlockAnal)**

Increment in the temperature during the previous time increment at the contact points on the master surface.

### **fieldSlv(nFields,nBlock)**

Current user-specified predefined field variables at the slave nodes (initial values at the beginning of the analysis and current values during the analysis).

### **fieldMst(nFields,nBlockAnal)**

Current user-specified predefined field variables at the contact points on the master surface (initial values at the beginning of the analysis and current values during the analysis).

### **dFieldSlv(nFields,nBlock)**

Increment in the user-specified predefined field variables during the previous time increment at the slave nodes.

### **dFieldMst(nFields,nBlockAnal)**

Increment in the user-specified predefined field variables during the previous time increment at the contact points on the master surface.

## **Additional information**

---

- “**VUINTERACTION**,” Section 4.1.36 of the Abaqus Verification Manual

## 1.2.17 VUMAT: User subroutine to define material behavior.

**Product:** Abaqus/Explicit

*WARNING: The use of this user subroutine generally requires considerable expertise. You are cautioned that the implementation of any realistic constitutive model requires extensive development and testing. Initial testing on a single-element model with prescribed traction loading is strongly recommended.*

*The component ordering of the symmetric and nonsymmetric tensors for the three-dimensional case using C3D8R elements is different from the ordering specified in “Three-dimensional solid element library,” Section 25.1.4 of the Abaqus Analysis User’s Manual, and the ordering used in Abaqus/Standard.*

---

## References

- “User-defined mechanical material behavior,” Section 23.8.1 of the Abaqus Analysis User’s Manual
- \*USER MATERIAL

---

## Overview

User subroutine **VUMAT**:

- is used to define the mechanical constitutive behavior of a material;
- will be called for blocks of material calculation points for which the material is defined in a user subroutine (“Material data definition,” Section 18.1.2 of the Abaqus Analysis User’s Manual);
- can use and update solution-dependent state variables;
- can use any field variables that are passed in; and
- can be used in an adiabatic analysis, provided you define both the inelastic heat fraction and the specific heat for the appropriate material definitions and you store the temperatures and integrate them as user-defined state variables.

---

## Component ordering in tensors

The component ordering depends upon whether the tensor is symmetric or nonsymmetric.

### Symmetric tensors

For symmetric tensors such as the stress and strain tensors, there are **ndir+nshr** components, and the component order is given as a natural permutation of the indices of the tensor. The direct components are first and then the indirect components, beginning with the 12-component. For example, a stress tensor contains **ndir** direct stress components and **nshr** shear stress components, which are passed in as

Component	2-D Case	3-D Case
1	$\sigma_{11}$	$\sigma_{11}$
2	$\sigma_{22}$	$\sigma_{22}$
3	$\sigma_{33}$	$\sigma_{33}$
4	$\sigma_{12}$	$\sigma_{12}$
5		$\sigma_{23}$
6		$\sigma_{31}$

The shear strain components in user subroutine **VUMAT** are stored as tensor components and not as engineering components; this is different from user subroutine **UMAT** in Abaqus/Standard, which uses engineering components.

### Nonsymmetric tensors

For nonsymmetric tensors there are **ndir+2\*nshr** components, and the component order is given as a natural permutation of the indices of the tensor. The direct components are first and then the indirect components, beginning with the 12-component. For example, the deformation gradient is passed as

Component	2-D Case	3-D Case
1	$F_{11}$	$F_{11}$
2	$F_{22}$	$F_{22}$
3	$F_{33}$	$F_{33}$
4	$F_{12}$	$F_{12}$
5	$F_{21}$	$F_{23}$
6		$F_{31}$
7		$F_{21}$
8		$F_{32}$
9		$F_{13}$

---

### Initial calculations and checks

In the data check phase of the analysis Abaqus/Explicit calls user subroutine **VUMAT** with a set of fictitious strains and a **totalTime** and **stepTime** both equal to 0.0. This is done as a check on your constitutive relation and to calculate the equivalent initial material properties, based upon which the initial elastic wave speeds are computed.

## Defining local orientations

---

All stresses, strains, stretches, and state variables are in the orientation of the local material axes. These local material axes form a basis system in which stress and strain components are stored. This represents a corotational coordinate system in which the basis system rotates with the material. If a user-specified coordinate system (“Orientations,” Section 2.2.5 of the Abaqus Analysis User’s Manual) is used, it defines the local material axes in the undeformed configuration.

## Special considerations for various element types

---

The use of user subroutine **VUMAT** requires special consideration for various element types.

### Shell and plane stress elements

You must define the stresses and internal state variables. In the case of shell or plane stress elements, **NDIR=3** and **NSHR=1**; you must define **strainInc(\*,3)**, the thickness strain increment. The internal energies can be defined if desired. If they are not defined, the energy balance provided by Abaqus/Explicit will not be meaningful.

### Shell elements

When **VUMAT** is used to define the material response of shell elements, Abaqus/Explicit cannot calculate a default value for the transverse shear stiffness of the element. Hence, you must define the element’s transverse shear stiffness. See “Shell section behavior,” Section 26.6.4 of the Abaqus Analysis User’s Manual, for guidelines on choosing this stiffness.

### Beam elements

For beam elements the stretch tensor and the deformation gradient tensor are not available. For beams in space you must define the thickness strains, **strainInc(\*,2)** and **strainInc(\*,3)**. **strainInc(\*,4)** is the shear strain associated with twist. Thickness stresses, **stressNew(\*,2)** and **stressNew(\*,3)**, are assumed to be zero, and any values you assign are ignored.

### Pipe elements

For pipe elements the stretch tensor and the deformation gradient tensor are not available. The axial strain, **strainInc(\*,1)**, and the shear strain, **strainInc(\*,4)**, associated with twist are provided along with the hoop stress, **stressNew(\*,2)**. The hoop stress is predefined based on your pipe internal and external pressure load definitions (PE, PI, HPE, HPI, PENU, and PINU), and it should not be modified here. The thickness stress, **stressNew(\*,3)**, is assumed to be zero and any value you assign is ignored. You must define the axial stress, **stressNew(\*,1)**, and the shear stress, **stressNew(\*,4)**. You must also define hoop strain, **strainInc(\*,2)**, and the pipe thickness strain, **strainInc(\*,3)**.

## Deformation gradient

---

The polar decomposition of the deformation gradient is written as  $\mathbf{F} = \mathbf{R} \cdot \mathbf{U} = \mathbf{V} \cdot \mathbf{R}$ , where  $\mathbf{U}$  and  $\mathbf{V}$  are the right and left symmetric stretch tensors, respectively. The constitutive model is defined in a corotational coordinate system in which the basis system rotates with the material. All stress and strain tensor quantities are defined with respect to the corotational basis system. The right stretch tensor,  $\mathbf{U}$ , is used. The relative spin tensor  $\mathbf{W} - \boldsymbol{\Omega}$  represents the spin (the antisymmetric part of the velocity gradient) defined with respect to the corotational basis system.

## Special considerations for hyperelasticity

Hyperelastic constitutive models in **VUMAT** should be defined in a corotational coordinate system in which the basis system rotates with the material. This is most effectively accomplished by formulating the hyperelastic constitutive model in terms of the stretch tensor,  $\mathbf{U}$ , instead of in terms of the deformation gradient,  $\mathbf{F} = \mathbf{R} \cdot \mathbf{U}$ . Using the deformation gradient can present some difficulties because the deformation gradient includes the rotation tensor and the resulting stresses would need to be rotated back to the corotational basis.

## Objective stress rates

---

The Green-Naghdi stress rate is used when the mechanical behavior of the material is defined using user subroutine **VUMAT**. The stress rate obtained with user subroutine **VUMAT** may differ from that obtained with a built-in Abaqus material model. For example, most material models used with solid (continuum) elements in Abaqus/Explicit employ the Jaumann stress rate. This difference in the formulation will cause significant differences in the results only if finite rotation of a material point is accompanied by finite shear. For a discussion of the objective stress rates used in Abaqus, see “Stress rates,” Section 1.5.3 of the Abaqus Theory Manual.

## Material point deletion

---

Material points that satisfy a user-defined failure criterion can be deleted from the model (see “User-defined mechanical material behavior,” Section 23.8.1 of the Abaqus Analysis User’s Manual). You must specify the state variable number controlling the element deletion flag when you allocate space for the solution-dependent state variables, as explained in “User-defined mechanical material behavior,” Section 23.8.1 of the Abaqus Analysis User’s Manual. The deletion state variable should be set to a value of one or zero in **VUMAT**. A value of one indicates that the material point is active, while a value of zero indicates that Abaqus/Explicit should delete the material point from the model by setting the stresses to zero. The structure of the block of material points passed to user subroutine **VUMAT** remains unchanged during the analysis; deleted material points are not removed from the block. Abaqus/Explicit will pass zero stresses and strain increments for all deleted material points. Once a material point has been flagged as deleted, it cannot be reactivated.

---

User subroutine interface

```

        subroutine vumat(
C Read only (unmodifiable) variables -
 1 nblock, ndir, nshr, nstatev, nfieldv, nprops, lanneal,
 2 stepTime, totalTime, dt, cmname, coordMp, charLength,
 3 props, density, strainInc, relSpinInc,
 4 tempOld, stretchOld, defgradOld, fieldOld,
 5 stressOld, stateOld, enerInternOld, enerInelasOld,
 6 tempNew, stretchNew, defgradNew, fieldNew,
C Write only (modifiable) variables -
 7 stressNew, stateNew, enerInternNew, enerInelasNew )
C
C           include 'vaba_param.inc'
C
      dimension props(nprops), density(nblock), coordMp(nblock,*),
 1 charLength(nblock), strainInc(nblock,ndir+nshr),
 2 relSpinInc(nblock,nshr), tempOld(nblock),
 3 stretchOld(nblock,ndir+nshr),
 4 defgradOld(nblock,ndir+nshr+nshr),
 5 fieldOld(nblock,nfieldv), stressOld(nblock,ndir+nshr),
 6 stateOld(nblock,nstatev), enerInternOld(nblock),
 7 enerInelasOld(nblock), tempNew(nblock),
 8 stretchNew(nblock,ndir+nshr),
 8 defgradNew(nblock,ndir+nshr+nshr),
 9 fieldNew(nblock,nfieldv),
 1 stressNew(nblock,ndir+nshr), stateNew(nblock,nstatev),
 2 enerInternNew(nblock), enerInelasNew(nblock),
C
C           character*80 cmname
C
      do 100 km = 1,nblock
      user coding
100 continue

      return
      end

```

---

**Variables to be defined****stressNew (nblock, ndir+nshr)**

Stress tensor at each material point at the end of the increment.

**stateNew (nblock, nstatev)**

State variables at each material point at the end of the increment. You define the size of this array by allocating space for it (see “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual, for more information).

---

**Variables that can be updated****enerInternNew (nblock)**

Internal energy per unit mass at each material point at the end of the increment.

**enerInelasNew (nblock)**

Dissipated inelastic energy per unit mass at each material point at the end of the increment.

---

**Variables passed in for information****nblock**

Number of material points to be processed in this call to **VUMAT**.

**ndir**

Number of direct components in a symmetric tensor.

**nshr**

Number of indirect components in a symmetric tensor.

**nstatev**

Number of user-defined state variables that are associated with this material type (you define this as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**lanneal**

Flag indicating whether the routine is being called during an annealing process. **lanneal**=0 indicates that the routine is being called during a normal mechanics increment. **lanneal**=1 indicates that this is an annealing process and you should re-initialize the internal state variables, **stateNew**, if necessary. Abaqus/Explicit will automatically set the stresses, stretches, and state to a value of zero during the annealing process.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime** - **stepTime**.

**dt**

Time increment size.

**cmname**

User-specified material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, you should not use “ABQ\_” as the leading string for **cmname**.

**coordMp (nblock, \*)**

Material point coordinates. It is the midplane material point for shell elements and the centroid for beam and pipe elements.

**charLength (nblock)**

Characteristic element length. This is a typical length of a line across an element for a first-order element; it is half of the same typical length for a second-order element. For beams, pipes, and trusses, it is a characteristic length along the element axis. For membranes and shells it is a characteristic length in the reference surface. For axisymmetric elements it is a characteristic length in the  $r-z$  plane only. For cohesive elements it is equal to the constitutive thickness.

**props (nprops)**

User-supplied material properties.

**density (nblock)**

Current density at the material points in the midstep configuration. This value may be inaccurate in problems where the volumetric strain increment is very small. If an accurate value of the density is required in such cases, the analysis should be run in double precision. This value of the density is not affected by mass scaling.

**strainInc (nblock, ndir+nshr)**

Strain increment tensor at each material point.

**relSpinInc (nblock, nshr)**

Incremental relative rotation vector at each material point defined in the corotational system. Defined as  $\Delta t(\mathbf{W} - \boldsymbol{\Omega})$ , where  $\mathbf{W}$  is the antisymmetric part of the velocity gradient,  $\mathbf{L}$ , and  $\boldsymbol{\Omega} = \dot{\mathbf{R}} \cdot \mathbf{R}^T$ . Stored in 3-D as (32, 13, 21) and in 2-D as (21).

**tempOld (nblock)**

Temperatures at each material point at the beginning of the increment.

**stretchOld (nblock, ndir+nshr)**

Stretch tensor,  $\mathbf{U}$ , at each material point at the beginning of the increment defined from the polar decomposition of the deformation gradient by  $\mathbf{F} = \mathbf{R} \cdot \mathbf{U}$ .

**defgradOld (nblock,ndir+2\*nshr)**

Deformation gradient tensor at each material point at the beginning of the increment. Stored in 3-D as  $(F_{11}, F_{22}, F_{33}, F_{12}, F_{23}, F_{31}, F_{21}, F_{32}, F_{13})$  and in 2-D as  $(F_{11}, F_{22}, F_{33}, F_{12}, F_{21})$ .

**fieldOld (nblock, nfieldv)**

Values of the user-defined field variables at each material point at the beginning of the increment.

**stressOld (nblock, ndir+nshr)**

Stress tensor at each material point at the beginning of the increment.

**stateOld (nblock, nstatev)**

State variables at each material point at the beginning of the increment.

**enerInternOld (nblock)**

Internal energy per unit mass at each material point at the beginning of the increment.

**enerInelasOld (nblock)**

Dissipated inelastic energy per unit mass at each material point at the beginning of the increment.

**tempNew (nblock)**

Temperatures at each material point at the end of the increment.

**stretchNew (nblock, ndir+nshr)**

Stretch tensor,  $\mathbf{U}$ , at each material point at the end of the increment defined from the polar decomposition of the deformation gradient by  $\mathbf{F} = \mathbf{R} \cdot \mathbf{U}$ .

**defgradNew (nblock,ndir+2\*nshr)**

Deformation gradient tensor at each material point at the end of the increment. Stored in 3-D as  $(F_{11}, F_{22}, F_{33}, F_{12}, F_{23}, F_{31}, F_{21}, F_{32}, F_{13})$  and in 2-D as  $(F_{11}, F_{22}, F_{33}, F_{12}, F_{21})$ .

**fieldNew (nblock, nfieldv)**

Values of the user-defined field variables at each material point at the end of the increment.

**Example: Using more than one user-defined material model**

---

To use more than one user-defined material model, the variable **cmname** can be tested for different material names inside user subroutine **VUMAT**, as illustrated below:

```
if (cmname(1:4) .eq. 'MAT1') then
    call VUMAT_MAT1(argument_list)
else if (cmname(1:4) .eq. 'MAT2') then
    call VUMAT_MAT2(argument_list)
end if
```

**VUMAT\_MAT1** and **VUMAT\_MAT2** are the actual user material subroutines containing the constitutive material models for each material **MAT1** and **MAT2**, respectively. Subroutine **VUMAT** merely acts as a directory here. The argument list can be the same as that used in subroutine **VUMAT**. The material names must be in uppercase characters since **cmname** is passed in as an uppercase character string.

### Example: Elastic/plastic material with kinematic hardening

---

As a simple example of the coding of subroutine **VUMAT**, consider the generalized plane strain case for an elastic/plastic material with kinematic hardening. The basic assumptions and definitions of the model are as follows.

Let  $\sigma$  be the current value of the stress, and define  $S$  to be the deviatoric part of the stress. The center of the yield surface in deviatoric stress space is given by the tensor  $\alpha$ , which has initial values of zero. The stress difference,  $\xi$ , is the stress measured from the center of the yield surface and is given by

$$\xi = S - \alpha .$$

The von Mises yield surface is defined as

$$f(\sigma) = \frac{1}{2}\xi : \xi - \frac{1}{3}\sigma_0^2,$$

where  $\sigma_0$  is the uniaxial equivalent yield stress. The von Mises yield surface is a cylinder in deviatoric stress space with a radius of

$$R = \sqrt{\frac{2}{3}}\sigma_0.$$

For the kinematic hardening model,  $R$  is a constant. The normal to the Mises yield surface can be written as

$$\mathbf{Q} = \sqrt{\frac{3}{2}} \frac{\xi}{\sigma_0} .$$

We decompose the strain rate into an elastic and plastic part using an additive decomposition:

$$\dot{\epsilon} = \dot{\epsilon}^{el} + \dot{\epsilon}^{pl} .$$

The plastic part of the strain rate is given by a normality condition

$$\dot{\epsilon}^{pl} = \dot{\gamma} \mathbf{Q},$$

where the scalar multiplier  $\dot{\gamma}$  must be determined. A scalar measure of equivalent plastic strain rate is defined by

$$\dot{\epsilon}^{pl} = \sqrt{\frac{2}{3}\dot{\epsilon}^{pl} : \dot{\epsilon}^{pl}} .$$

The stress rate is assumed to be purely due to the elastic part of the strain rate and is expressed in terms of Hooke's law by

$$\dot{\sigma} = \lambda \text{trace}(\dot{\epsilon}^{el}) \mathbf{I} + 2\mu \dot{\epsilon}^{el},$$

where  $\lambda$  and  $2\mu$  are the Lamé constants for the material.

The evolution law for  $\alpha$  is given as

$$\dot{\alpha} = \frac{2}{3} \dot{\gamma} H \mathbf{Q},$$

where  $H$  is the slope of the uniaxial yield stress versus plastic strain curve.

During active plastic loading the stress must remain on the yield surface, so that

$$\sqrt{\mathbf{Q} : \mathbf{Q}} = 1.$$

The equivalent plastic strain rate is related to  $\dot{\gamma}$  by

$$\dot{\epsilon}^{pl} = \sqrt{\frac{2}{3}} \dot{\gamma}.$$

The kinematic hardening constitutive model is integrated in a rate form as follows. A trial elastic stress is computed as

$$\sigma_{new}^{trial} = \sigma_{old} + \lambda \text{trace}(\Delta\epsilon) \mathbf{I} + 2\mu \Delta\epsilon,$$

where the subscripts *old* and *new* refer to the beginning and end of the increment, respectively. If the trial stress does not exceed the yield stress, the new stress is set equal to the trial stress. If the yield stress is exceeded, plasticity occurs in the increment. We then write the incremental analogs of the rate equations as

$$\sigma_{new} = \sigma_{new}^{trial} - 2\mu \Delta\epsilon^{pl} = \sigma_{new}^{trial} - 2\mu \Delta\gamma \mathbf{Q},$$

$$\alpha_{new} = \alpha_{old} + \frac{2}{3} H \Delta\gamma \mathbf{Q},$$

$$\bar{\epsilon}_{new}^{pl} = \bar{\epsilon}_{old}^{pl} + \sqrt{\frac{2}{3}} \Delta\gamma,$$

where

$$\Delta\gamma = \dot{\gamma} \Delta t.$$

From the definition of the normal to the yield surface at the end of the increment,  $\mathbf{Q}$ ,

$$\alpha_{new} + \sqrt{\frac{2}{3}}\sigma_0 \mathbf{Q} = \mathbf{S}_{new}.$$

This can be expanded using the incremental equations as

$$\alpha_{old} + \frac{2}{3}H\Delta\gamma\mathbf{Q} + \sqrt{\frac{2}{3}}\sigma_0 \mathbf{Q} = \mathbf{S}_{new}^{trial} - \Delta\gamma 2\mu\mathbf{Q}.$$

Taking the tensor product of this equation with  $\mathbf{Q}$ , using the yield condition at the end of the increment, and solving for  $\Delta\gamma$ :

$$\Delta\gamma = \frac{1}{2\mu(1+H/3\mu)} \left( (\xi_{new}^{trial} : \xi_{new}^{trial})^{1/2} - \sqrt{\frac{2}{3}}\sigma_0 \right).$$

The value for  $\Delta\gamma$  is used in the incremental equations to determine  $\sigma_{new}$ ,  $\alpha_{new}$ , and  $\bar{\epsilon}_{new}^{pl}$ .

This algorithm is often referred to as an elastic predictor, radial return algorithm because the correction to the trial stress under the active plastic loading condition returns the stress state to the yield surface along the direction defined by the vector from the center of the yield surface to the elastic trial stress. The subroutine would be coded as follows:

```

subroutine vumat(
C Read only -
  1 nblock, ndir, nshr, nstatev, nfieldv, nprops, lanneal,
  2 stepTime, totalTime, dt, cmname, coordMp, charLength,
  3 props, density, strainInc, relSpinInc,
  4 tempOld, stretchOld, defgradOld, fieldOld,
  3 stressOld, stateOld, enerInternOld, enerInelasOld,
  6 tempNew, stretchNew, defgradNew, fieldNew,
C Write only -
  5 stressNew, stateNew, enerInternNew, enerInelasNew )
C
      include 'vaba_param.inc'
C
C J2 Mises Plasticity with kinematic hardening for plane
C strain case.
C Elastic predictor, radial corrector algorithm.
C
C The state variables are stored as:
C     STATE(*,1) = back stress component 11
C     STATE(*,2) = back stress component 22
C     STATE(*,3) = back stress component 33
C     STATE(*,4) = back stress component 12
C     STATE(*,5) = equivalent plastic strain

```

```

C
C
C All arrays dimensioned by (*) are not used in this algorithm
      dimension props(nprops), density(nblock),
1   coordMp(nblock,*),
2   charLength(*), strainInc(nblock,ndir+nshr),
3   relSpinInc(*), tempOld(*),
4   stretchOld(*), defgradOld(*),
5   fieldOld(*), stressOld(nblock,ndir+nshr),
6   stateOld(nblock,nstatev), enerInternOld(nblock),
7   enerInelasOld(nblock), tempNew(*),
8   stretchNew(*), defgradNew(*), fieldNew(*),
9   stressNew(nblock,ndir+nshr), stateNew(nblock,nstatev),
1   enerInternNew(nblock), enerInelasNew(nblock)

C
      character*80 cmname
C
      parameter( zero = 0., one = 1., two = 2., three = 3.,
1   third = one/three, half = .5, twoThirds = two/three,
2   threeHalfs = 1.5 )
C
      e      = props(1)
      xnu   = props(2)
      yield = props(3)
      hard  = props(4)
C
      twomu = e / ( one + xnu )
      thremu = threeHalfs * twomu
      sixmu = three * twomu
      alamda = twomu * ( e - twomu ) / ( sixmu - two * e )
      term   = one / ( twomu * ( one + hard/thremu ) )
      con1   = sqrt( twoThirds )
C
      do 100 i = 1,nblock
C
C Trial stress
      trace = strainInc(i,1) + strainInc(i,2) + strainInc(i,3)
      sig1 = stressOld(i,1) + alamda*trace + twomu*strainInc(i,1)
      sig2 = stressOld(i,2) + alamda*trace + twomu*strainInc(i,2)
      sig3 = stressOld(i,3) + alamda*trace + twomu*strainInc(i,3)
      sig4 = stressOld(i,4)           + twomu*strainInc(i,4)
C

```

```

C Trial stress measured from the back stress
    s1 = sig1 - stateOld(i,1)
    s2 = sig2 - stateOld(i,2)
    s3 = sig3 - stateOld(i,3)
    s4 = sig4 - stateOld(i,4)
C
C Deviatoric part of trial stress measured from the back stress
    smean = third * ( s1 + s2 + s3 )
    ds1 = s1 - smean
    ds2 = s2 - smean
    ds3 = s3 - smean
C
C Magnitude of the deviatoric trial stress difference
    dsmag = sqrt( ds1**2 + ds2**2 + ds3**2 + 2.*s4**2 )
C
C Check for yield by determining the factor for plasticity,
C zero for elastic, one for yield
    radius = con1 * yield
    facyld = zero
    if( dsmag - radius .ge. zero ) facyld = one
C
C Add a protective addition factor to prevent a divide by zero
C when dsmag is zero. If dsmag is zero, we will not have exceeded
C the yield stress and facyld will be zero.
    dsmag = dsmag + ( one - facyld )
C
C Calculated increment in gamma (this explicitly includes the
C time step)
    diff = dsmag - radius
    dgamma = facyld * term * diff
C
C Update equivalent plastic strain
    deqps = con1 * dgamma
    stateNew(i,5) = stateOld(i,5) + deqps
C
C Divide dgamma by dsmag so that the deviatoric stresses are
C explicitly converted to tensors of unit magnitude in the
C following calculations
    dgamma = dgamma / dsmag
C
C Update back stress
    factor = hard * dgamma * twoThirds

```

```

stateNew(i,1) = stateOld(i,1) + factor * ds1
stateNew(i,2) = stateOld(i,2) + factor * ds2
stateNew(i,3) = stateOld(i,3) + factor * ds3
stateNew(i,4) = stateOld(i,4) + factor * s4
C
C Update the stress
    factor = twomu * dgamma
    stressNew(i,1) = sig1 - factor * ds1
    stressNew(i,2) = sig2 - factor * ds2
    stressNew(i,3) = sig3 - factor * ds3
    stressNew(i,4) = sig4 - factor * s4
C
C Update the specific internal energy -
    stressPower = half * (
        1      ( stressOld(i,1)+stressNew(i,1) )*strainInc(i,1)
        1      +      ( stressOld(i,2)+stressNew(i,2) )*strainInc(i,2)
        1      +      ( stressOld(i,3)+stressNew(i,3) )*strainInc(i,3)
        1      + two*( stressOld(i,4)+stressNew(i,4) )*strainInc(i,4) )
C
        enerInternNew(i) = enerInternOld(i)
        1      + stressPower / density(i)
C
C Update the dissipated inelastic specific energy -
    plasticWorkInc = dgamma * half * (
        1      ( stressOld(i,1)+stressNew(i,1) )*ds1
        1      +      ( stressOld(i,2)+stressNew(i,2) )*ds2
        1      +      ( stressOld(i,3)+stressNew(i,3) )*ds3
        1      + two*( stressOld(i,4)+stressNew(i,4) )*s4 )
    enerInelasNew(i) = enerInelasOld(i)
    1      + plasticWorkInc / density(i)
100 continue
C
    return
end

```

## 1.2.18 VUSDFLD: User subroutine to redefine field variables at a material point.

**Product:** Abaqus/Explicit

### References

---

- “Obtaining material point information in an Abaqus/Standard analysis,” Section 2.1.6
- “Material data definition,” Section 18.1.2 of the Abaqus Analysis User’s Manual
- \*USER DEFINED FIELD
- “Damage and failure of a laminated composite plate,” Section 1.1.14 of the Abaqus Example Problems Manual
- “**VUSDFLD**,” Section 4.1.38 of the Abaqus Verification Manual

### Overview

---

User subroutine **VUSDFLD**:

- allows the redefinition of field variables at a material point as functions of time or of any of the available material point quantities listed in “Available output variable keys” in “Obtaining material point information in an Abaqus/Explicit analysis,” Section 2.1.7;
- can be used to introduce solution-dependent material properties since such properties can be easily defined as functions of field variables;
- will be called at all material points of elements for which the material definition includes user-defined field variables;
- can call utility routine **VGETVRM** to access material point data; and
- can use and update solution-dependent state variables.

### Explicit solution dependence

---

Since this routine provides access to material point quantities only at the start of the increment, the material properties for a given increment are not influenced by the results obtained during the increment. Hence, the accuracy of the results depends on the size of the time increment. However, in most situations this is not a concern for explicit dynamic analysis because the stable time increment is usually sufficiently small to ensure good accuracy.

### Defining field variables

---

Before user subroutine **VUSDFLD** is called, the values of the field variables at the material point are calculated by interpolation from the values defined at the nodes. Any changes to the field variables in the user subroutine are local to the material point: the nodal field variables retain the values defined as initial conditions or predefined field variables or the values defined in user subroutine **VUFIELD**. The values of the field variables defined in this routine are used to calculate values of material properties that

are defined to depend on field variables and are passed into other user subroutines that are called at the material point, such as the following:

- **VUANISOHYPER\_INV**
- **VUANISOHYPER\_STRAIN**
- **VUHARD**
- **VUMAT**
- **VUTRS**
- **VUVISCOSITY**

Output of the user-defined field variables at the material points can be obtained with the element integration point output variable FV (see “Abaqus/Explicit output variable identifiers,” Section 4.2.2 of the Abaqus Analysis User’s Manual).

## State variables

---

Since the redefinition of field variables in **VUSDFLD** is local to the current increment (field variables are restored to the values interpolated from the nodal values at the start of each increment), any history dependence required to update material properties by using this subroutine must be introduced with user-defined state variables.

The state variables can be updated in **VUSDFLD** and then passed into other user subroutines that can be called at this material point, such as those listed above. The number of such state variables can be specified as shown in the example at the end of this section (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

## Accessing material point data

---

The values of the material point quantities at the start of the increment can be accessed through the utility routine **VGETVRM** described in “Obtaining material point information in an Abaqus/Explicit analysis,” Section 2.1.7. The values of the material point quantities are obtained by calling **VGETVRM** with the appropriate output variable keys.

## Component ordering in symmetric tensors

---

For symmetric tensors such as the stress and strain tensors there are **ndir+nshr** components, and the component order is given as a natural permutation of the indices of the tensor. The direct components are first and then the indirect components, beginning with the 12-component. For example, a stress tensor contains **ndir** direct stress components and **nshr** shear stress components, which are returned as:

Component	2-D Case	3-D Case
1	$\sigma_{11}$	$\sigma_{11}$
2	$\sigma_{22}$	$\sigma_{22}$
3	$\sigma_{33}$	$\sigma_{33}$

Component	2-D Case	3-D Case
4	$\sigma_{12}$	$\sigma_{12}$
5		$\sigma_{23}$
6		$\sigma_{31}$

The shear strain components in user subroutine **VUSDFLD** are stored as tensor components and not as engineering components; unlike user subroutine **USDFLD** in Abaqus/Standard, which uses engineering components.

### User subroutine interface

---

```

subroutine vusdfld(
c Read only variables -
 1  nblock, nstatev, nfieldv, nprops, ndir, nshr,
 2  jElem, kIntPt, kLayer, kSecPt,
 3  stepTime, totalTime, dt, cmname,
 4  coordMp, direct, T, charLength, props,
 5  stateOld,
c Write only variables -
 6  stateNew, field )
c
      include 'vaba_param.inc'
c
      dimension jElem(nblock), coordMp(nblock,*),
 1          direct(nblock,3,3), T(nblock,3,3),
 2          charLength(nblock), props(nprops),
 3          stateOld(nblock,nstatev),
 4          stateNew(nblock,nstatev),
 5          field(nblock,nfieldv)
      character*80 cmname
c
c Local arrays from vgetvrm are dimensioned to
c maximum block size (maxblk)
c
      parameter( nrData=6 )
      character*3 cData(maxblk*nrData)
      dimension rData(maxblk*nrData), jData(maxblk*nrData)
c
      do 100 k = 1, nblock

      user coding to define field(nblock,nfieldv)
      and, if necessary, stateNew(nblock,nstatev)

```

```
100 continue
c
      return
end
```

---

**Variable to be defined****field(nblock, nfieldv)**

An array containing the field variables at the material points. These are passed in with the values interpolated from the nodes at the end of the current increment, as specified with initial condition definitions, predefined field variable definitions, or user subroutine **VUFIELD**. The updated values are used to calculate the values of material properties that are defined to depend on field variables and are passed into other user subroutines that are called at the material points.

---

**Variable that can be updated****stateNew(nblock, nstatev)**

An array containing the solution-dependent state variables at the material points. In all cases **stateNew** can be updated in this subroutine, and the updated values are passed into other user subroutines that are called at the material points. The number of state variables associated with this material point is defined as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

---

**Variables passed in for information****nblock**

Number of material points to be processed in this call to **VUSDFLD**.

**nstatev**

Number of user-defined state variables that are associated with this material type (you define this as described in “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**ndir**

Number of direct components in a symmetric tensor.

**nshr**

Number of indirect components in a symmetric tensor.

**jElem**

Array of element numbers.

**kIntPt**

Integration point number.

**kLayer**

Layer number (for composite shells).

**kSecPt**

Section point number within the current layer.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**dt**

Time increment size.

**cmname**

User-specified material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, you should not use “ABQ\_” as the leading string for **cmname**.

**coordMp (nblock, \*)**

Material point coordinates. It is the midplane material point for shell elements and the centroid for beam elements.

**direct (nblock, 3, 3)**

An array containing the direction cosines of the material directions in terms of the global basis directions. For material point k, **direct(k,1,1)**, **direct(k,2,1)**, **direct(k,3,1)** give the (1, 2, 3) components of the first material direction; **direct(k,1,2)**, **direct(k,2,2)**, **direct(k,3,2)** give the second material direction, etc. For shell and membrane elements, the first two directions are in the plane of the element and the third direction is the normal. This information is not available for beam elements.

**T(nblock, 3, 3)**

An array containing the direction cosines of the material orientation components relative to the element basis directions. For material point k, this is the orientation that defines the material directions (**direct**) in terms of the element basis directions. For continuum elements **T** and **direct** are identical. For shell and membrane elements **T(k,1,1)** =  $\cos \theta$ , **T(k,1,2)** =  $-\sin \theta$ , **T(k,2,1)** =  $\sin \theta$ , **T(k,2,2)** =  $\cos \theta$ , **T(k,3,3)** = 1.0 and all other components are zero, where  $\theta$  is the

counterclockwise rotation around the normal vector that defines the orientation. If no orientation is used,  $\mathbf{T}$  is an identity matrix. Orientation is not available for beam elements.

**charLength (nblock)**

Characteristic element length. This is a typical length of a line across an element for a first-order element; it is half of the same typical length for a second-order element. For beams and trusses, it is a characteristic length along the element axis. For membranes and shells it is a characteristic length in the reference surface. For axisymmetric elements it is a characteristic length in the  $r-z$  plane only. For cohesive elements it is equal to the constitutive thickness.

**props (nprops)**

User-supplied material properties.

**stateOld (nblock, nstatev)**

State variables at each material point at the beginning of the increment.

---

**Example: Damaged elasticity model**

Included below is an example of user subroutine **VUSDFLD**. In this example a truss element is loaded in tension. A damaged elasticity model is introduced: the modulus decreases as a function of the maximum tensile strain that occurred during the loading history. The maximum tensile strain is stored as a solution-dependent state variable (see “Defining solution-dependent field variables” in “Predefined fields,” Section 30.6.1 of the Abaqus Analysis User’s Manual).

**Input file**

```
*HEADING
  Damaged elasticity model with user subroutine vusdfld
*ELEMENT, TYPE=T2D2, ELSET=ONE
  1, 1, 2
*NODE, NSET=NALL
  1, 0., 0.
  2, 10., 0.
*SOLID SECTION, ELSET=ONE, MATERIAL=ELASTIC
  1.
*MATERIAL, NAME=ELASTIC
*ELASTIC, DEPENDENCIES=1
** Table of modulus values decreasing as a function
** of field variable 1.
  2000., 0.3, 0., 0.00
  1500., 0.3, 0., 0.01
  1200., 0.3, 0., 0.02
  1000., 0.3, 0., 0.04
*DENSITY
  1.0e-6
```

```

*USER DEFINED FIELD
*DEPVAR
1
1,EPSSMAX,"Maximum strain value"
*BOUNDARY
1, 1, 2
2, 2
*AMPLITUDE, NAME=LOAD1
0.0, 0.0, 1.0, 1.0
*AMPLITUDE, NAME=LOAD2
0.0, 0.0, 2.0, 1.0
*AMPLITUDE, NAME=UNLOAD
0.0, 1.0, 1.0, 0.0
*STEP, NLGEOM=NO
*DYNAMIC, EXPLICIT
, 1.0
*CLOAD, AMPLITUDE=LOAD1
2, 1, 20.
*OUTPUT, FIELD, VARIABLE=PRESELECT
*OUTPUT, HISTORY, VARIABLE=PRESELECT
*ELEMENT OUTPUT, ELSET=ONE
S, E, SDV
*NODE OUTPUT, NSET=NALL
RF, CF, U
*END STEP
*STEP, NLGEOM=NO
*DYNAMIC, EXPLICIT
, 1.0
*CLOAD, AMPLITUDE=UNLOAD
2, 1, 20.
*END STEP
*STEP, NLGEOM=NO
*DYNAMIC, EXPLICIT
, 2.0
*CLOAD, AMPLITUDE=LOAD2
2, 1, 40.
*END STEP

```

#### User subroutine

```

      subroutine vusdfld(
c Read only -
      *   nblock, nstatev, nfieldv, nprops, ndir, nshr,

```

```

*      jElem, kIntPt, kLayer, kSecPt,
*      stepTime, totalTime, dt, cmname,
*      coordMp, direct, T, charLength, props,
*      stateOld,
c Write only -
*      stateNew, field )
c
include 'vaba_param.inc'
c
dimension jElem(nblock), coordMp(nblock,*),
*           direct(nblock,3,3), T(nblock,3,3),
*           charLength(nblock), props(nprops),
*           stateOld(nblock,nstatev),
*           stateNew(nblock,nstatev),
*           field(nblock,nfieldv)
character*80 cmname
c
c Local arrays from vgetvrm are dimensioned to
c maximum block size (maxblk)
c
parameter( nrData=6 )
character*3 cData(maxblk*nrData)
dimension rData(maxblk*nrData), jData(maxblk*nrData)
c
jStatus = 1
call vgetvrm( 'LE', rData, jData, cData, jStatus )
c
if( jStatus .ne. 0 ) then
    call xplb_abqerr(-2,'Utility routine VGETVRM '//
*          'failed to get variable.',0,zero,' ')
    call xplb_exit
end if
c
call setField( nblock, nstatev, nfieldv, nrData,
*   rData, stateOld, stateNew, field)
c
return
end
subroutine setField( nblock, nstatev, nfieldv, nrData,
*   strain, stateOld, stateNew, field )
c
include 'vaba_param.inc'

```

```
c
dimension stateOld(nblock,nstatev) ,
*      stateNew(nblock,nstatev) ,
*      field(nblock,nfieldv), strain(nblock,nrData)
c
do k = 1, nblock
c
c   Absolute value of current strain:
      eps = abs( strain(k,1) )
c
c   Maximum value of strain up to this point in time:
      epsmax = stateOld(k,1)
c
c   Use the maximum strain as a field variable
      field(k,1) = max( eps, epsmax )
c
c   Store the maximum strain as a solution dependent state
      stateNew(k,1) = field(k,1)
c
end do
c
return
end
```



## 1.2.19 VUTRS: User subroutine to define a reduced time shift function for a viscoelastic material.

**Product:** Abaqus/Explicit

### References

---

- “Time domain viscoelasticity,” Section 19.7.1 of the Abaqus Analysis User’s Manual
- \*TRS
- \*VISCOELASTIC
- “Transient thermal loading of a viscoelastic slab,” Section 3.1.2 of the Abaqus Benchmarks Manual

### Overview

---

User subroutine **VUTRS**:

- can be used to define a temperature-time shift for a time domain viscoelastic analysis;
- will be called for all material points of elements for which a user-defined shift function is specified to define the time-temperature correspondence as part of the viscoelastic material definition;
- can use and update solution-dependent state variables; and
- can have incoming field variables redefined by user subroutine **VUSDFLD**.

### User subroutine interface

---

```

subroutine vutrs(
c Read only variables -
  1 nblock, nstatev, nfieldv, nprops,
  2 stepTime, totalTime, dt,
  3 cmname, props, density, coordMp,
  4 tempOld, fieldOld, stateOld,
  5 tempNew, fieldNew,
c Write only variables -
  6 shift, stateNew )
c
  include 'vaba_param.inc'
c
  dimension props(nprops), density(nblock), coordMp(nblock,*),
  1 tempOld(nblock),tempNew(nblock),
  2 fieldOld(nblock,nfieldv), fieldNew(nblock,nfieldv),
  3 stateOld(nblock,nstatev), stateNew(nblock,nstatev),
  4 shift(nblock,2)

```

```

c
character*80 cmname
c
do 100 k=1, nblock
  user coding to define shift(k,1) and shift(k,2)
100 continue
c
return
end

```

---

**Variable to be defined****shift(nblock,2)**

Array that defines the shift function,  $A$  ( $A > 0$ ), at the material points. For material point  $k$ , **shift(k,1)** defines the shift function at the beginning of the increment, and **shift(k,2)** defines the shift function at the end of the increment. Abaqus/Explicit will apply an averaging scheme to these values that assumes that the natural logarithm of the shift function can be approximated by a linear function over the increment.

If either **shift(k,1)** or **shift(k,2)** is less than or equal to zero, no time shift will be applied.

---

**Variable that can be updated****stateNew(nblock,nstatev)**

Array containing the solution-dependent state variables at the material points. This array will be passed in containing the values of these variables at the start of the increment unless they are updated in user subroutine **VUSDFLD**, in which case the updated values are passed in. If any of the solution-dependent state variables are being used in conjunction with the viscoelastic behavior, they must be updated in this subroutine to their values at the end of the increment.

---

**Variables passed in for information****nblock**

Number of material points to be processed in this call to **VUTRS**.

**nstatev**

Number of user-defined state variables that are associated with this material type (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**stepTime**

Value of time since the step began.

**totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**dt**

Time increment size.

**cmname**

Material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, “ABQ\_” should not be used as the leading string for **cmname**.

**props (nprops)**

User-supplied material properties.

**density (nblock)**

Current density at the material points in the midstep configuration. This value may be inaccurate in problems where the volumetric strain increment is very small. If an accurate value of the density is required in such cases, the analysis should be run in double precision. This value of the density is not affected by mass scaling.

**coordMp (nblock, \*)**

Material point coordinates. It is the midplane material point for shell elements and the centroid for beam elements.

**tempOld (nblock)**

Temperatures at each material point at the beginning of the increment.

**fieldOld (nblock, nfieldv)**

Values of the user-defined field variables at each material point at the beginning of the increment.

**stateOld (nblock, nstatev)**

State variables at each material point at the beginning of the increment.

**tempNew (nblock)**

Temperatures at each material point at the end of the increment.

**fieldNew (nblock, nfieldv)**

Values of the user-defined field variables at each material point at the end of the increment.



## 1.2.20 UVISCOSITY: User subroutine to define the shear viscosity for equation of state models.

**Product:** Abaqus/Explicit

### References

---

- “Equation of state,” Section 22.2.1 of the Abaqus Analysis User’s Manual
- \*EOS
- \*VISCOSITY
- “**UVISCOSITY**,” Section 4.1.39 of the Abaqus Verification Manual

### Overview

---

User subroutine **UVISCOSITY**:

- is called at all material points of elements with an equation of state for which the material definition includes user-defined viscous shear behavior;
- can be used to define a material’s isotropic viscous behavior;
- can use and update solution-dependent state variables; and
- can be used in conjunction with user subroutine **VUSDFLD** to redefine any field variables before they are passed in.

### User subroutine interface

---

```

        subroutine uvviscosity(
C Read only -
        *      nblock,
        *      jElem, kIntPt, kLayer, kSecPt,
        *      stepTime, totalTime, dt, cmname,
        *      nstatev, nfieldv, nprops,
        *      props, tempOld, tempNew, fieldOld, fieldNew,
        *      stateOld,
        *      shrRate,
C Write only -
        *      viscosity,
        *      stateNew )
C
        include 'vaba_param.inc'
C
        dimension props(nprops), tempOld(nblock), tempNew(nblock),

```

## VUVISCOSITY

```
1  fieldOld(nblock,nfieldv), fieldNew(nblock,nfieldv),
2  stateOld(nblock,nstatev), eqps(nblock), eqpsRate(nblock),
3  viscosity(nblock),
4  stateNew(nblock,nstatev), jElem(nblock)
C
      character*80 cmname
C
      do 100 km = 1,nblock
         user coding
100 continue
C
      return
end
```

### Variables to be defined

---

#### **viscosity(nblock)**

Array containing the viscosity at the material points. (Units of  $\text{FL}^{-2}\text{T}$ .)

#### **stateNew(nblock,nstatev)**

Array containing the state variables at the material points at the end of the increment. The allocation of this array is described in “Solution-dependent state variables” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual.

### Variables passed in for information

---

#### **nblock**

Number of material points to be processed in this call to **VUVISCOSITY**.

#### **jElem(nblock)**

Array of element numbers.

#### **kIntPt**

Integration point number.

#### **kLayer**

Layer number (for composite shells).

#### **kSecPt**

Section point number within the current layer.

#### **stepTime**

Value of time since the step began.

#### **totalTime**

Value of total time. The time at the beginning of the step is given by **totalTime-stepTime**.

**dt**

Time increment size.

**cmname**

Material name, left justified. It is passed in as an uppercase character string. Some internal material models are given names starting with the “ABQ\_” character string. To avoid conflict, “ABQ\_” should not be used as the leading string for **cmname**.

**nstatev**

Number of user-defined state variables that are associated with this material type (see “Allocating space” in “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual).

**nfieldv**

Number of user-defined external field variables.

**nprops**

User-specified number of user-defined material properties.

**tempOld(nblock)**

Temperatures at the material points at the beginning of the increment.

**tempNew(nblock)**

Temperatures at the material points at the end of the increment.

**fieldOld(nblock,nfieldv)**

Values of the user-defined field variables at the material points at the beginning of the increment.

**fieldNew(nblock,nfieldv)**

Values of the user-defined field variables at the material points at the end of the increment.

**stateOld(nblock,nstatev)**

State variables at the material points at the beginning of the increment.

**shrRate(nblock)**

Equivalent shear strain rate,  $\dot{\gamma}$ , at the material points.

**Example: Cross viscosity model**

---

As a simple example of the coding of subroutine **VUVISCOSITY**, consider the Cross viscosity model. The Cross model is commonly used when it is necessary to describe the low shear rate behavior of the viscosity. The viscosity is expressed as

$$\eta = \frac{\eta_0}{1 + (\lambda\dot{\gamma})^{1-n}},$$

## VUVISCOSITY

where  $\eta_0$  is the Newtonian viscosity,  $n$  is the flow index in the power law regime, and  $\lambda$  is a constant with units of time, such that  $1/\lambda$  corresponds to the critical shear rate at which the fluid changes from Newtonian to power law behavior.

The subroutine would be coded as follows:

```
      subroutine vuviscosity (
C Read only -
      *      nblock,
      *      jElem, kIntPt, kLayer, kSecPt,
      *      stepTime, totalTime, dt, cmname,
      *      nstatev, nfieldv, nprops,
      *      props, tempOld, tempNew, fieldOld, fieldNew,
      *      stateOld,
      *      shrRate,
C Write only -
      *      viscosity,
      *      stateNew )
C
      include 'vaba_param.inc'
C
      dimension props(nprops),
      *      tempOld(nblock),
      *      fieldOld(nblock,nfieldv),
      *      stateOld(nblock,nstatev),
      *      shrRate(nblock),
      *      tempNew(nblock),
      *      fieldNew(nblock,nfieldv),
      *      viscosity(nblock),
      *      stateNew(nblock,nstatev)
C
      character*80 cmname
C
      parameter ( one = 1.d0 )
C
C      Cross viscosity
C
      eta0      = props(1)
      rlambda   = props(2)
      rn        = props(3)
C
      do k = 1, nblock
         viscosity(k) = eta0/(one+(rlambda*shrRate(k))** (one-rn))
      end do
```

```
c
return
end
```



## 2. Utility Routines

---

- “Utility routines,” Section 2.1



## 2.1 Utility routines

- “Obtaining Abaqus environment variables,” Section 2.1.1
- “Obtaining the Abaqus job name,” Section 2.1.2
- “Obtaining the Abaqus output directory name,” Section 2.1.3
- “Obtaining parallel processes information,” Section 2.1.4
- “Obtaining part information,” Section 2.1.5
- “Obtaining material point information in an Abaqus/Standard analysis,” Section 2.1.6
- “Obtaining material point information in an Abaqus/Explicit analysis,” Section 2.1.7
- “Obtaining material point information averaged at a node,” Section 2.1.8
- “Obtaining node point information,” Section 2.1.9
- “Obtaining node to element connectivity,” Section 2.1.10
- “Obtaining stress invariants, principal stress/strain values and directions, and rotating tensors in an Abaqus/Standard analysis,” Section 2.1.11
- “Obtaining principal stress/strain values and directions in an Abaqus/Explicit analysis,” Section 2.1.12
- “Obtaining wave kinematic data in an Abaqus/Aqua analysis,” Section 2.1.13
- “Printing messages to the message or status file,” Section 2.1.14
- “Terminating an analysis,” Section 2.1.15
- “Obtaining sensor information,” Section 2.1.16
- “Accessing Abaqus materials,” Section 2.1.17
- “Accessing Abaqus thermal materials,” Section 2.1.18



## 2.1.1 OBTAINING Abaqus ENVIRONMENT VARIABLES

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- “Using the Abaqus environment settings,” Section 3.3.1 of the Abaqus Analysis User’s Manual
- “**UWAVE** and **UEXTERNALDB**,” Section 4.1.27 of the Abaqus Verification Manual

### Overview

---

Utility routines **GETENVVAR** and **VGETENVVAR** can be called from any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, to obtain the value of an environment variable.

### Interface

---

```
character*256 ENVVAR
...
CALL GETENVVAR( 'ENVVARNAMEx', ENVVAR, LENVVAR )
CALL VGETENVVAR( 'ENVVARNAMEx', ENVVAR, LENVVAR )
...
```

#### Variable to be provided to the utility routine

---

##### **ENVVARNAMEx**

Environment variable name.

#### Variables returned from the utility routine

---

##### **ENVVAR**

Character string to receive the value of the environment variable.

##### **LENVVAR**

Length of the character string **ENVVAR**.



## 2.1.2 OBTAINING THE Abaqus JOB NAME

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- “**UWAVE** and **UEXTERNALDB**,” Section 4.1.27 of the Abaqus Verification Manual

### Overview

---

Utility routines **GETJOBNAME** and **VGETJOBNAME** can be called from any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, to obtain the name of the current job.

### Interface

---

```
character*256 JOBNAME  
...  
CALL GETJOBNAME( JOBNAME, LENJOBNAME )  
CALL VGETJOBNAME( JOBNAME, LENJOBNAME )  
...
```

### Variables returned from the utility routine

---

#### **JOBNAME**

Character string to receive the value of the job name.

#### **LENJOBNAME**

Length of the character string **JOBNAME**.



## 2.1.3 OBTAINING THE Abaqus OUTPUT DIRECTORY NAME

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- “**UWAVE** and **UEXTERNALDB**,” Section 4.1.27 of the Abaqus Verification Manual

### Overview

---

Utility routines **GETOUTDIR** and **VGETOUTDIR** can be called from any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, to obtain the output directory of the current job.

### Interface

---

```
character*256 OUTDIR
...
CALL GETOUTDIR( OUTDIR, LENOUTDIR )
CALL VGETOUTDIR( OUTDIR, LENOUTDIR )
...
```

### Variables returned from the utility routine

---

#### OUTDIR

Character string to receive the value of the output directory name.

#### LENOUTDIR

Length of the character string **OUTDIR**.



## 2.1.4 OBTAINING PARALLEL PROCESSES INFORMATION

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “Parallel execution: overview,” Section 3.5.1 of the Abaqus Analysis User’s Manual
- “**VUSDFLD**,” Section 4.1.38 of the Abaqus Verification Manual

### Overview

---

Utility routines **GETNUMCPUS** and **GETRANK** can be called from any Abaqus/Standard user subroutine. **GETNUMCPUS** returns the number of MPI processes, and **GETRANK** returns the rank of the MPI process from which the function is called. For example, in a hybrid MPI and thread parallel execution scheme, multiple threads may all return the rank of their parent MPI process (see “Parallel execution in Abaqus/Standard,” Section 3.5.2 of the Abaqus Analysis User’s Manual).

Utility routines **VGETNUMCPUS** and **VGETRANK** can be called from any Abaqus/Explicit user subroutine in a domain-parallel run. **VGETNUMCPUS** provides the number of processes used for the parallel run, and **VGETRANK** provides the individual process rank (see “Parallel execution in Abaqus/Explicit,” Section 3.5.3 of the Abaqus Analysis User’s Manual).

### GETNUMCPUS and VGETNUMCPUS (obtain the number of processes)

---

#### Interface

```
CALL GETNUMCPUS( NUMPROCESSES )
CALL VGETNUMCPUS( NUMPROCESSES )
...

```

#### Variable returned from the utility routine

##### NUMPROCESSES

Number of processes specified for the analysis.

### GETRANK and VGETRANK (obtain the process number)

---

#### Interface

```
CALL GETRANK( KPROCESSNUM )
CALL VGETRANK( KPROCESSNUM )
...

```

## OBTAINING PROCESS INFORMATION

### **Variable returned from the utility routine**

#### **KPROCESSNUM**

Process number or rank. A process number is either zero or a positive integer.

## 2.1.5 OBTAINING PART INFORMATION

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- “Defining an assembly,” Section 2.9.1 of the Abaqus Analysis User’s Manual
- “Pure bending of a cylinder: CAXA elements,” Section 1.3.33 of the Abaqus Verification Manual

### Overview

---

Utility routines **GETPARTINFO** and **VGETPARTINFO** can be called from any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, to retrieve the part instance name and original node or element number corresponding to an internal node or element number. Utility routines **GETINTERNAL** and **VGETINTERNAL** can be called from any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, to retrieve the internal node or element number corresponding to a part instance name and original node or element number. The part file (*jobname.prt*) must be available. The expense of calling these routines is not trivial, so minimal use of them is recommended.

### GETPARTINFO and VGETPARTINFO (obtain part instance information given global node/element number)

---

#### Interface

```
CHARACTER*80 CPNAME
...
CALL GETPARTINFO(INTNUM, JTYP, CPNAME, LOCNUM, JRCD)
or
CALL VGETPARTINFO(INTNUM, JTYP, CPNAME, LOCNUM, JRCD)
```

#### Variables to be provided to the utility routine

##### **INTNUM**

The internal (global) node or element number to be looked up.

##### **JTYP**

An integer flag indicating whether **INTNUM** is a node or element number. Set **JTYP=0** to look up a node number, and set **JTYP=1** to look up an element number.

## OBTAINING PART INFORMATION

### Variables returned from the utility routine

#### **CPNAME**

The name of the part instance that contains **INTNUM**. An empty part instance name indicates that the node or element is at the assembly level and is not included in any part instance.

#### **LOCMNUM**

The part-local node or element number corresponding to **INTNUM**.

#### **JRCD**

Return code (0–no error, 1–error).

---

### **GETINTERNAL and VGETINTERNAL (obtain global node/element number given part instance information )**

---

### Interface

```
CHARACTER*80 CPNAME  
...  
CALL GETINTERNAL(CPNAME, LOCMNUM, JTYP, INTNUM, JRCD)  
or  
CALL VGETINTERNAL(CPNAME, LOCMNUM, JTYP, INTNUM, JRCD)
```

### Variables to be provided to the utility routine

#### **CPNAME**

The name of the part instance that contains **LOCMNUM**.

#### **LOCMNUM**

The part-local node or element number to be looked up.

#### **JTYP**

An integer flag indicating whether **LOCMNUM** is a node or element number. Set **JTYP**=0 to look up a node number, and set **JTYP**=1 to look up an element number.

### Variables returned from the utility routine

#### **INTNUM**

The internal (global) node or element number corresponding to **LOCMNUM** in part instance **CPNAME**.

#### **JRCD**

Return code (0–no error, 1–error).

## 2.1.6 OBTAINING MATERIAL POINT INFORMATION IN AN Abaqus/Standard ANALYSIS

**Product:** Abaqus/Standard

### References

---

- “UVARM,” Section 1.1.50
- “USDFLD,” Section 1.1.46
- “UDMGINI,” Section 1.1.23
- “Damage and failure of a laminated composite plate,” Section 1.1.14 of the Abaqus Example Problems Manual
- “**USDFLD**,” Section 4.1.24 of the Abaqus Verification Manual
- “**UVARM**,” Section 4.1.26 of the Abaqus Verification Manual

### Overview

---

Utility routine **GETVRM** can be called from either user subroutine **UVARM**, **UDMGINI**, or **USDFLD** to access material integration point information.

### Interface

---

```
DIMENSION ARRAY(15), JARRAY(15)
CHARACTER*3 FLGRAY(15)
...
CALL GETVRM('VAR',ARRAY,JARRAY,FLGRAY,JRCD,JMAC,JMATYP,MATLAYO,
LACCFLA)
```

### Variables to be provided to the utility routine

---

#### **VAR**

Output variable key from the table in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual. The applicable keys are listed in the output table as being available for results file output at the element integration points; e.g., S for stress.

#### **JMAC**

Variable that must be passed into the **GETVRM** utility routine. The calling user subroutine provides this variable.

#### **JMATYP**

Variable that must be passed into the **GETVRM** utility routine. The calling user subroutine provides this variable.

### **MATLAYO**

Variable that must be passed into the **GETVRM** utility routine. The calling user subroutine provides this variable.

### **LACCFLA**

Variable that must be passed into the **GETVRM** utility routine. The calling user subroutine provides this variable.

## **Variables returned from the utility routine**

---

### **ARRAY**

Real array containing individual components of the output variable.

### **JARRAY**

Integer array containing individual components of the output variable.

### **FLGRAY**

Character array containing flags corresponding to the individual components. Flags will contain either YES, NO, or N/A (not applicable).

### **JRCD**

Return code (0 – no error, 1 – output request error or all components of output request are zero).

## **Available output variable keys**

---

Only output variable keys that are valid for results file output are available for use with **GETVRM**. In general, if a key corresponds to a collective output variable, rather than an individual component, it can be used with **GETVRM**. For example, S for the stress tensor can be used, whereas any individual component of stress, say S11, cannot be used. The collective output variable keys are distinguished from their individual components by the fact that they have a bullet (•) in the .fil column in the tables in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual. Output variable keys that cannot be used with **GETVRM** are listed later in this section.

You will be returned **ARRAY**, **JARRAY**, and **FLGRAY**, which correspond to the real-valued components, integer-valued components, and the flags associated with the request **VAR**, respectively. If any array component is not applicable for a given request, its value will be returned as the initialized value: 0.0 in **ARRAY**, 0 in **JARRAY**, and N/A in **FLGRAY**. The error flag **JRCD=1** is returned from **GETVRM** any time a request key is not recognized, the request is not valid (such as requesting transverse shear stress for a shell element that uses thin shell theory), or all of the output components requested are zero; otherwise, **JRCD=0**.

## **Ordering of returned components**

---

The components for a request are written as follows. Single index components (and requests without components) are returned in positions 1, 2, 3, etc. Double index components are returned in the

order 11, 22, 33, 12, 13, 23 for symmetric tensors, followed by 21, 31, 32 for unsymmetric tensors (deformation gradient). Thus, the stresses for a plane stress element are returned as **ARRAY(1)**=S11, **ARRAY(2)**=S22, **ARRAY(3)**=0.0, and **ARRAY(4)**=S12. Three values are always returned for principal value requests, the minimum value first and maximum value third, regardless of the dimensionality of the analysis.

The description of the output variable (see “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual) determines which components are retrieved with **GETVRM**.

### **Analysis time for which values are returned**

---

When a material point quantity is requested with utility routine **GETVRM**, the time in the increment at which the values are returned will depend upon which user subroutine calls it. **GETVRM** returns values at the end of the current increment to user subroutine **UVARM**, whereas it returns values at the beginning of the current increment to user subroutine **USDFLD**.

### **Equilibrium state for values returned**

---

User subroutine **UVARM** may call **GETVRM** multiple times for each increment, as Abaqus/Standard iterates to a converged solution. Values returned from **GETVRM** calls preceding the final iteration for the increment will not represent the converged solution.

### **Example**

---

To illustrate the use of **GETVRM**, if the identifier PEQC is specified for use with a jointed material, **ARRAY** will be returned with the individual equivalent plastic strain components PEQC1, PEQC2, PEQC3, and PEQC4. Since there are no integer output variables associated with this identifier, **JARRAY** will be returned with default values of 0. The **FLGRAY** array will contain either YES or NO flags indicating whether each component is actively yielding. If the identifier PE is specified for a material with plasticity, **ARRAY** will be returned with plastic strain components PE11, PE22, PE33, PE12, PE13, PE23, the equivalent plastic strain PEEQ, and the plastic strain magnitude PEMAG. Since there are no integer values associated with this request, **JARRAY** will be 0. The **FLGRAY** array will have N/A for the first six components, either YES or NO in the seventh component (corresponding to PEEQ) indicating whether the material is currently yielding, and N/A in the eighth component. If the identifier HFL is specified, **ARRAY** will be returned with the magnitude HFLM and the components HFL1, HFL2, and HFL3 as described in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual.

### **Accessing state-dependent variables**

---

If **GETVRM** is used to access state-dependent variables (output variable key SDV) and more than 15 state-dependent variables have been defined in the analysis, the dimension statement for **ARRAY** and **JARRAY** must be changed so that these arrays are dimensioned to the maximum number of state-dependent variables.

---

**Unsupported element types, procedures and output variable keys**

Since this capability pertains to material point quantities, it cannot be used for most of the element types that do not require a material definition. The following element types are, therefore, not supported:

- DASHPOTx
- SPRINGx
- CONNxDx
- FRAMExD
- JOINTC
- JOINTxD
- DRAGxD
- PSIxx
- ITSxxx
- MASS
- ROTARYI
- all acoustic elements
- all contact elements
- all hydrostatic fluid elements

If used with user subroutine **UVARM**, this capability is not available for linear perturbation procedures (“General and linear perturbation procedures,” Section 6.1.2 of the Abaqus Analysis User’s Manual):

- static linear perturbation analysis (“Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual),
- “Eigenvalue buckling prediction,” Section 6.2.3 of the Abaqus Analysis User’s Manual,
- “Natural frequency extraction,” Section 6.3.5 of the Abaqus Analysis User’s Manual,
- “Transient modal dynamic analysis,” Section 6.3.7 of the Abaqus Analysis User’s Manual,
- “Mode-based steady-state dynamic analysis,” Section 6.3.8 of the Abaqus Analysis User’s Manual,
- “Direct-solution steady-state dynamic analysis,” Section 6.3.4 of the Abaqus Analysis User’s Manual,
- “Subspace-based steady-state dynamic analysis,” Section 6.3.9 of the Abaqus Analysis User’s Manual,
- “Response spectrum analysis,” Section 6.3.10 of the Abaqus Analysis User’s Manual, and
- “Random response analysis,” Section 6.3.11 of the Abaqus Analysis User’s Manual.

The following output variable keys are not available for use with **GETVRM**:

- SVOL
- TSHR
- CTSHR
- COORD

## 2.1.7 OBTAINING MATERIAL POINT INFORMATION IN AN Abaqus/Explicit ANALYSIS

**Product:** Abaqus/Explicit

### References

---

- “VUSDFLD,” Section 1.2.18
- “Damage and failure of a laminated composite plate,” Section 1.1.14 of the Abaqus Example Problems Manual

### Overview

---

Utility routine **VGETVRM** can be called from **VUSDFLD** to access selected output variables at the material points for the current block of elements being processed by **VUSDFLD**.

### Interface

---

```
include 'vaba_param.inc'
parameter( nrData=6 )
character*3 cData(maxblk*nrData)
dimension rData(maxblk*nrData), jData(maxblk*nrData)
...
call vgetvrm( 'VAR', rData, jData, cData, jStatus )
```

### Variable to be provided to the utility routine

---

#### **VAR**

Output variable key. The available material point variables are given in “Available output variable keys,” below.

### Variables returned from the utility routine

---

#### **rData**

Real array containing individual components of the output variable.

#### **jData**

Integer array containing individual components of the output variable.

#### **cData**

Character array containing flags corresponding to the individual components. Flags will either be YES, NO, or N/A (not applicable).

**jStatus**

Return code (0: output request successful; 1: output request not available).

**Available output variable keys**

---

The following output variable keys are currently supported:

- S: All stress components.
- LE: All logarithmic strain components.
- THE: All thermal strain components.
- PE: All plastic strain components.
- PEEQ: Equivalent plastic strain.
- PEEQT: Equivalent plastic strain in uniaxial tension, defined as  $\int \dot{\varepsilon}_t^{pl} dt$ .
- PEQC: All equivalent plastic strains for models that have more than one yield/failure surface.
- ALPHA: All total kinematic hardening shift tensor components.
- TEMP: Temperature.
- EVF: Volume fraction (Eulerian elements only).

A requested output variable must be valid for the material model for the request to be successful.

The returned arrays **rData**, **jData**, and **cData** correspond to the real-valued variable, integer-valued variable, and string variable, respectively, that can be associated with the request output variable key **VAR**. If any output variable component is not applicable for a given request, its value will be returned as the initialized value: 0.0 in **rData**, 0 in **jData**, and N/A in **cData**. Currently the association of the integer-valued variable and the string variable with the request output variable is not supported. The error flag **jStatus** is returned with a value of 1 if a request key is not recognized, not valid, or not supported; otherwise, **jStatus** is returned with a value of 0.

**Component ordering in symmetric tensors**

---

For symmetric tensors such as the stress and strain tensors, there are **ndir+nshr** components, where **ndir** and **nshr** are the number of direct and shear components, respectively, that are passed into user subroutine **VUSDFLD**. The component order is given as a natural permutation of the indices of the tensor. The direct components are first and then the indirect components, beginning with the 12-component. For example, a stress tensor contains **ndir** direct stress components and **nshr** shear stress components, which are returned as:

Component	2-D Case	3-D Case
1	$\sigma_{11}$	$\sigma_{11}$
2	$\sigma_{22}$	$\sigma_{22}$
3	$\sigma_{33}$	$\sigma_{33}$
4	$\sigma_{12}$	$\sigma_{12}$

Component	2-D Case	3-D Case
5		$\sigma_{23}$
6		$\sigma_{31}$

The shear strain components returned from utility subroutine **VGETVRM** are tensor components and not engineering components.

#### **Analysis time for which values are returned**

---

Utility subroutine **VGETVRM** returns values of the requested variable that correspond to the beginning of the current increment.

#### **Example**

---

To illustrate the use of **VGETVRM**, if the identifier PE is specified for a material with plasticity, **rData** will be returned with plastic strain components PE11, PE22, PE33, PE12, PE23, and PE13. **jData** will be 0 and **cData** array will have N/A for all components.

#### **Unsupported element types, procedures, and output variable keys**

---

Since this capability pertains to material point quantities, it cannot be used for most of the element types that do not require a material definition. The following element types are, therefore, not supported:

- DASHPOTx
- SPRINGx
- CONNxDx
- MASS
- ROTARYI
- all acoustic elements



## 2.1.8 OBTAINING MATERIAL POINT INFORMATION AVERAGED AT A NODE

**Product:** Abaqus/Standard

### References

---

- “UMESHMOTION,” Section 1.1.39
- “Erosion of material (sand production) in an oil wellbore,” Section 1.1.22 of the Abaqus Example Problems Manual

### Overview

---

Utility routine **GETVRMAVGATNODE** can be called from user subroutine **UMESHMOTION** to access material integration point information averaged at a node. The results variables available from **GETVRMAVGATNODE** are nearly the same as those available from **GETVRM**; the exceptions follow from the restriction that, since it will average results, **GETVRMAVGATNODE** will operate only on real-valued results. Results values represented as integers or as flags are not available.

### Interface

---

```
DIMENSION ARRAY(15) , JELEMLIST(NELEMS)
...
CALL GETVRMAVGATNODE (NODE , JTYP , 'VAR' , ARRAY , JRCD , JELEMLIST , NELEMS ,
JMATYP , JGVBLOCK)
```

### Variables to be provided to the utility routine

---

#### NODE

Node number.

#### JTYP

An integer flag indicating how the material point information is averaged. Set **JTYP**=0 to extrapolate results, using element shape functions, and to average results at the node. Set **JTYP**=1 to perform a volume-weighted average of results.

#### VAR

Output variable key from the table in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual. The applicable keys are listed in the output table as being available for results file output at the element integration points; e.g., S for stress. One exception is the integration point coordinates variable COORD, which cannot be passed into the utility routine; you should use utility routine **GETVRN** instead to obtain nodal coordinates.

**JELEMLIST**

Array of element numbers for elements connected to **NODE** for which you want material point quantities considered in the average result. Results from each element in the list that contain the node will be extrapolated to that node and averaged. **JELEMLIST** can be obtained from utility routine **GETNODETOELEMCONN**.

**NELEMS**

Length of **JELEMLIST**.

**JGVBLOCK**

Variable that must be passed into the **GETVRMAVGATNODE** utility routine. This variable is available in user subroutine **UMESHMOTION** for this purpose.

**JMATYP**

Variable that must be passed into the **GETVRMAVGATNODE** utility routine. This variable is available in user subroutine **UMESHMOTION** for this purpose.

**Variables returned from the utility routine**

---

**ARRAY**

Real array containing individual components of the output variable.

**JRCD**

Return code (0 – no error, 1 – output request error or all components of output request are zero).

**Available output variable keys**

---

Only output variable keys that are valid for results file output are available for use with **GETVRMAVGATNODE**. In general, if a key corresponds to a collective output variable, rather than an individual component, it can be used with **GETVRMAVGATNODE**. For example, S for the stress tensor can be used, whereas any individual component of stress, say S11, cannot be used. The collective output variable keys are distinguished from their individual components by the fact that they have a bullet (•) in the .fil column in the tables in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual. Output variable keys that cannot be used with **GETVRMAVGATNODE** are listed later in this section.

You will be returned **ARRAY** with components associated with the request **VAR**. If any array component is not applicable for a given request, its value will be returned as the initialized value: 0.0 in **ARRAY**. The error flag **JRCD=1** is returned from **GETVRMAVGATNODE** any time a request key is not recognized, the request is not valid, or all of the output components requested are zero; otherwise, **JRCD=0**.

## Ordering of returned components

---

The components for a request are written as follows. Single index components (and requests without components) are returned in positions 1, 2, 3, etc. Double index components are returned in the order 11, 22, 33, 12, 13, 23 for symmetric tensors, followed by 21, 31, 32 for unsymmetric tensors (deformation gradient). Thus, the stresses for a plane stress element are returned as **ARRAY (1)**=S11, **ARRAY (2)**=S22, **ARRAY (3)**=0.0, and **ARRAY (4)**=S12. Three values are always returned for principal value requests, the minimum value first and the maximum value third, regardless of the dimensionality of the analysis.

The description of the output variable (see “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual) determines which components are retrieved with **GETVRMAVGATNODE**.

## Analysis time for which values are returned

---

**GETVRMAVGATNODE** returns values at the end of the current increment to user subroutine **UMESHMOTION**.

## Accessing state-dependent variables

---

If **GETVRMAVGATNODE** is used to access solution-dependent state variables (output variable key SDV) and more than 15 solution-dependent state variables have been defined in the analysis, the dimension statement for **ARRAY** must be changed so that these arrays are dimensioned to the maximum number of solution-dependent state variables.

## Unsupported element types and output variable keys

---

Since this capability pertains to material point quantities, it cannot be used for most of the element types that do not require a material definition. The following element types are, therefore, not supported:

- DASHPOTx
- SPRINGx
- CONNxDx
- FRAMExD
- JOINTC
- JOINTxD
- DRAGxD
- PSIxx
- ITSxxx
- MASS
- ROTARYI

## MATERIAL POINT INFORMATION AT A NODE

- all acoustic elements
- all hydrostatic fluid elements

The following output variable keys are not available for use with **GETVRMAVGATNODE**:

- SVOL
- TSHR
- CTSHR

### **Example: Obtaining plastic strain results**

---

To illustrate the use of **GETVRMAVGATNODE**, consider a case where the identifier PE is specified and **JELEMLIST** lists four three-dimensional elements, two of which have plastic yield behavior defined and two of which do not. **ARRAY** will be returned with the individual plastic strain components PE11, PE22, PE33, PE12, PE13, and PE23; the equivalent plastic strain PEEQ; and the plastic strain magnitude PEMAG. The result returned in **ARRAY** will be an average reflecting extrapolations of plastic strain results to **NODE** from only the two elements that have plastic yield behavior defined.

### **Example: Obtaining contact results**

---

A second illustration is relevant to the modeling of wear with **UMESHMOTION**. Consider a case where **JELEMLIST** is obtained from **GETNODETOELEMCONN** and where the identifier CSTRESS is specified. If **NODE** is associated with a contact pair slave surface, **JELEMLIST** will contain the internal element identifier for the contact element associated with the slave node pairing. **ARRAY** will be returned with the individual contact stress components CPRESS, CSHEAR1, and CSHEAR2. Similarly, if CDISP is specified, **ARRAY** will be returned with the individual contact stress components CDISP, CSLIP1, and CSLIP2.

## 2.1.9 OBTAINING NODE POINT INFORMATION

**Product:** Abaqus/Standard

### References

---

- “UMESHMOTION,” Section 1.1.39
- “Erosion of material (sand production) in an oil wellbore,” Section 1.1.22 of the Abaqus Example Problems Manual

### Overview

---

Utility routine **GETVRN** can be called from user subroutine **UMESHMOTION** to access node point information.

### Interface

---

```
DIMENSION ARRAY(15),JGVBLOCK(*)  
...  
CALL GETVRN(NODE,'VAR',ARRAY,JRCD,JGVBLOCK,LTRN)
```

### Variables to be provided to the utility routine

---

#### NODE

Node number.

#### VAR

Output variable key from the table in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual. The applicable keys are listed in the output table as being available for results file output at nodes; e.g., U for displacement.

#### JGVBLOCK

Variable that must be passed into the **GETVRN** utility routine. The variable is available in user subroutine **UMESHMOTION** for this purpose.

#### LTRN

Variable indicating the coordinate system the nodal quantity is to be returned in. A value of 0 specifies that the results are to be returned in the global coordinate system, regardless of any transformation applied at the node. A value of 1 specifies that the results are to be returned in the local transformed system.

---

### Variables returned from the utility routine

---

**ARRAY**

Real array containing individual components of the output variable.

**JRCD**

Return code (0 – no error, 1 – output request error or all components of output request are zero).

---

### Available output variable keys

---

Only nodal output variable keys that are valid for results file output in the current step are available for use with **GETVRN**. In general, if a key corresponds to a collective output variable, rather than an individual component, it can be used with **GETVRN**. For example, U for displacement can be used, whereas any individual component of displacement, say U1, cannot be used. The collective output variable keys are distinguished from their individual components by the fact that they have a bullet (•) in the .fil column in the tables in “Abaqus/Standard output variable identifiers,” Section 4.2.1 of the Abaqus Analysis User’s Manual.

You will be returned **ARRAY**, which corresponds to the real-valued components associated with the request **VAR**. If any array component is not applicable for a given request, its value will be returned as the initialized value: 0.0. The error flag **JRCD=1** is returned from **GETVRN** any time a request key is not recognized, the request is not valid (such as requesting pore pressure for a node not associated with a pore pressure or acoustic element, or requesting a variable not available for the current procedure), or all of the output components requested are zero; otherwise, **JRCD=0**.

---

### Ordering of returned components

---

The components for a vector request are returned in positions 1, 2, 3, etc.

---

### Analysis time for which values are returned

---

**GETVRN** returns values at the end of the current increment.

## 2.1.10 OBTAINING NODE TO ELEMENT CONNECTIVITY

**Product:** Abaqus/Standard

### References

---

- “UMESHMOTION,” Section 1.1.39
- “Obtaining material point information averaged at a node,” Section 2.1.8
- “Erosion of material (sand production) in an oil wellbore,” Section 1.1.22 of the Abaqus Example Problems Manual

### Overview

---

Utility routine **GETNODETOELEMCONN** can be called from user subroutine **UMESHMOTION** to retrieve a list of elements connected to a specified node.

### Interface

---

```
DIMENSION JELEMLIST(*), JELEMTYPE(*), JGVBLOCK(*)
...
CALL GETNODETOELEMCONN(NODE, NELEMS, JELEMLIST, JELEMTYPE,
JRCD, JGVBLOCK)
```

### Variables to be provided to the utility routine

---

#### NODE

User node number.

#### NELEMS

Maximum allowable length of **JELEMLIST**.

#### JGVBLOCK

Variable that must be passed into the **GETNODETOELEMCONN** utility routine. This variable is available in user subroutine **UMESHMOTION** for this purpose.

### Variables returned from the utility routine

---

#### JELEMLIST

Array of element numbers for elements connected to **NODE**. The list will contain elements only in adaptive mesh domains active in the step as well as any contact elements associated with the domain.

## OBTAINING NODE CONNECTIVITY

### **JELEMTYPE**

Array describing the element types for each element entry in **JELEMLIST**.

**JELEMTYPE** entries:

- 1 indicates a solid element.
- 2 indicates a contact element.

### **NELEMS**

Actual length of **JELEMLIST**.

### **JRCD**

Return code (0 – no error, 1 – output request error or all components of output request are zero).

## 2.1.11 OBTAINING STRESS INVARIANTS, PRINCIPAL STRESS/STRAIN VALUES AND DIRECTIONS, AND ROTATING TENSORS IN AN Abaqus/Standard ANALYSIS

**Product:** Abaqus/Standard

### References

---

- “UMAT,” Section 1.1.37
- “Calculation of principal stresses and strains and their directions: FPRIN,” Section 13.1.3 of the Abaqus Example Problems Manual

### Overview

---

For Abaqus/Standard user subroutines that store stress and strain components according to the convention presented in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual, a number of utility routines are available for calculating stress invariants, principal stress/strain values, and principal stress/strain directions from the relevant tensors. There is also a utility routine available for transforming tensors to a new basis. These routines are most commonly called from user subroutine **UMAT**.

The following utility subroutines are available in Abaqus/Standard to perform tensor operations:

- **SINV** (calculate stress invariants)
- **SPRINC** (calculate principal values)
- **SPRIND** (calculate principal values and directions)
- **ROTSIG** (rotate a tensor)

These utility subroutines are described below in alphabetical order.

### **SINV (calculate stress invariants)**

---

#### Interface

```
CALL SINV(STRESS,SINV1,SINV2,NDI,NSHR)
```

#### Variables to be provided to the utility routine

##### **STRESS**

A stress tensor.

##### **NDI**

Number of direct components.

##### **NSHR**

Number of shear components.

## INVARIANTS AND PRINCIPAL VALUES

### Variables returned from the utility routine

#### SINV1

First invariant.

$$\text{SINV1} = \frac{1}{3} \text{trace } \sigma,$$

where  $\sigma$  is the stress tensor.

#### SINV2

Second invariant.

$$\text{SINV2} = \sqrt{\frac{3}{2} \mathbf{S} : \mathbf{S}},$$

where  $\mathbf{S}$  is the deviatoric stress tensor, defined as

$$\mathbf{S} = \sigma - \frac{1}{3} \text{trace } \sigma \mathbf{I}.$$

---

### SPRINC (calculate principal values)

#### Interface

```
CALL SPRINC (S, PS, LSTR, NDI, NSHR)
```

### Variables to be provided to the utility routine

#### S

Stress or strain tensor.

#### LSTR

An identifier. **LSTR**=1 indicates that **S** contains stresses; **LSTR**=2 indicates that **S** contains strains.

#### NDI

Number of direct components.

#### NSHR

Number of shear components.

### Variables returned from the utility routine

#### PS(I), I=1,2,3

The three principal values.

**SPRIND (calculate principal values and directions)****Interface**

```
CALL SPRIND (S, PS, AN, LSTR, NDI, NSHR)
```

**Variables to be provided to the utility routine****S**

A stress or a strain tensor.

**LSTR**

An identifier. **LSTR**=1 indicates that **S** contains stresses; **LSTR**=2 indicates that **S** contains strains.

**NDI**

Number of direct components.

**NSHR**

Number of shear components.

**Variables returned from the utility routine****PS (I) , I=1,2,3**

The three principal values.

**AN (K1, I) , I=1,2,3**

The direction cosines of the principal directions corresponding to **PS (K1)**.

**ROTSIG (rotate a tensor)****Interface**

```
CALL ROTSIG (S, R, SPRIME, LSTR, NDI, NSHR)
```

**Variables to be provided to the utility routine****S**

A stress or strain tensor.

**NDI**

Number of direct components.

**NSHR**

Number of shear components.

## INVARIANTS AND PRINCIPAL VALUES

**R**

Rotation matrix.

**LSTR**

An identifier. **LSTR** = 1 indicates **S** contains stresses; **LSTR** = 2 indicates **S** contains strains.

### Variable returned from the utility routine

**SPRIME**

The rotated stress or strain tensor.

### Typical usage

In user subroutine **UMAT** it is often necessary to rotate tensors during a finite-strain analysis. The matrix **DROT** that is passed into **UMAT** represents the incremental rotation of the material basis system in which the stress and strain are stored. For an elastic-plastic material that hardens isotropically, the elastic and plastic strain tensors must be rotated to account for the evolution of the material directions. In this case **S** is the elastic or plastic strain tensor and **R** is the incremental rotation **DROT**.

## 2.1.12 OBTAINING PRINCIPAL STRESS/STRAIN VALUES AND DIRECTIONS IN AN Abaqus/Explicit ANALYSIS

**Product:** Abaqus/Explicit

### Reference

---

- “VUMAT,” Section 1.2.17

### Overview

---

For Abaqus/Explicit user subroutines that store stress and strain components according to the convention presented in “Conventions,” Section 1.2.2 of the Abaqus Analysis User’s Manual, a number of utility routines are available for calculating principal stress/strain values and principal stress/strain directions from the relevant tensors. These routines are most commonly called from user subroutine **VUMAT**.

The following utility subroutines are available in Abaqus/Explicit to perform tensor operations:

- **VSPRINC** (calculate principal values)
- **VSPRIND** (calculate principal values and directions)

These utility subroutines are described below.

#### **VSPRINC (calculate principal values)**

---

##### Interface

```
call vsprinc( nblock, s, eigVal, ndir, nshr )
```

##### Variables to be provided to the utility routine

**s (nblock,ndir+nshr)**

Stress or strain symmetric tensor.

**nblock**

Number of material points to be processed in this call to **VSPRINC**.

**ndir**

Number of direct components in the symmetric tensor.

**nshr**

Number of shear components in the symmetric tensor.

**Variables returned from the utility routine**

**eigVal (nblock, I) , I=1,2,3**

The three principal values.

**VSPRIND (calculate principal values and directions)**

---

**Interface**

```
call vsprind( nblock, s, eigVal, eigVec, ndir, nshr )
```

**Variables to be provided to the utility routine**

**s (nblock,ndir+nshr)**

Stress or strain symmetric tensor.

**nblock**

Number of material points to be processed in this call to **VSPRIND**.

**ndir**

Number of direct components in the symmetric tensor.

**nshr**

Number of shear components in the symmetric tensor.

**Variables returned from the utility routine**

**eigVal (nblock, I) , I=1,2,3**

The three principal values.

**eigVec (nblock, I, K1) , I=1,2,3**

The direction cosines of the principal directions corresponding to **eigVal (K1)**.

## 2.1.13 OBTAINING WAVE KINEMATIC DATA IN AN Abaqus/Aqua ANALYSIS

**Product:** Abaqus/Aqua

### References

---

- “UEL,” Section 1.1.24
- “UWAVE,” Section 1.1.51
- “Abaqus/Aqua analysis,” Section 6.11.1 of the Abaqus Analysis User’s Manual
- \*AQUA
- “UEL,” Section 4.1.14 of the Abaqus Verification Manual

### Overview

---

Utility routines **GETWAVE**, **GETWAVEVEL**, **GETWINDVEL**, and **GETCURRVEL** are provided to access the fluid kinematic data for an Abaqus/Aqua analysis. These routines can be used only from within user subroutine **UEL**.

### GETWAVE (get wave kinematics)

---

#### Interface

```

PARAMETER (MWCOMP=number of wave components)
DIMENSION WAMP (MWCOMP) , WPERD (MWCOMP) , WXLAMB (MWCOMP) ,
1 WPHI (MWCOMP) , WOFF (3) , WANG (2 , MWCOMP)
...
CALL GETWAVE (MWCOMP , NWCOMP , WAMP , WPERD , WXLAMB , WPHI , WOFF , WANG ,
1 ELEVB , ELEVS , JTYPE , JRCD)

```

#### Variables returned from the utility routine

##### **NWCOMP**

Number of wave components (always 1 for Stokes wave theory).

##### **WAMP**

Array containing the amplitude of the wave components.

##### **WPERD**

Array containing the period of the wave components.

##### **WXLAMB**

Array containing the wavelength of the wave components.

**WPHI**

Array containing the phase angle of the wave components.

**WOFF**

Used only for gridded wave data (**JWTYPE**=2), when **WOFF** gives the position of the origin of the gridded coordinate system with respect to the global system.

**WANG (2,\*)**

For Stokes fifth-order wave theory **WANG (1,1)** and **WANG (2,1)** are the direction cosines of wave travel. For Airy wave theory **WANG (1,K1)** and **WANG (2,K1)** are the direction cosines of the direction of travel of the **K1**th wave. For gridded wave data **WANG (1,1)** and **WANG (2,1)** are the direction cosines of the wave data grid. In all cases these direction cosines are with respect to the global coordinate system.

**ELEVB**

User-defined elevation of the seabed.

**ELEVS**

User-defined elevation of the still water surface.

**JWTYPE**

Integer flag indicating the wave type, as follows:

<b>JWTYPE=0</b>	Airy wave theory
<b>JWTYPE=1</b>	Stokes fifth-order wave theory
<b>JWTYPE=2</b>	Wave data obtained from gridded values

**JRCD**

The error flag **JRCD** is returned from **GETWAVE** as 0 if all the wave kinematic data are read correctly and as -1 if an error occurred (for instance, **NWCOMP** is greater than **MWCOMP**).

---

**GETWAVEVEL, GETWINDVEL, and GETCURRVEL (get wave, wind, and current velocities)**

---

**Interface**

```
CALL GETWAVEVEL (NDIM, X, V, A, LERROR, NOEL, XINTERMED)
CALL GETWINDVEL (NDIM, X, V, NOEL, XINTERMED)
CALL GETCURRVEL (NDIM, X, V, NOEL, XINTERMED)
```

## Variables to be provided to the utility routine

### **NDIM**

Dimensionality of the element. It should be set to 2 for two-dimensional cases (for example, beams in a plane) and 3 for three-dimensional cases (for example, beams in space).

### **X(1..NDIM)**

Global coordinates of the point.

## Variables returned from the utility routine

### **V(1..NDIM)**

Velocity components in the global coordinate system.

### **A(1..NDIM)**

Wave acceleration components in the global coordinate system. This variable is returned by **GETWAVEVEL** only.

### **LERROR**

For gridded wave data **LERROR** is returned as 0 if the current point is within the grid or above the crest; it is returned as 1 if the point is outside the bounds of the grid. For Airy and Stokes waves **LERROR** is always returned as 0. If **LERROR** is returned as 1, the global coordinates of the nearest grid point are returned in **X**. **LERROR** is returned by **GETWAVEVEL** only.

### **NOEL**

Element number.

### **XINTERMED (NDIM)**

An array containing the intermediate configuration coordinates of the load integration point. For nonstochastic analysis this array is not used. In a stochastic analysis the wave field is based upon this configuration. Additional details are found in “UWAVE,” Section 1.1.51.



## 2.1.14 PRINTING MESSAGES TO THE MESSAGE OR STATUS FILE

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- “**UWAVE** and **UEXTERNALDB**,” Section 4.1.27 of the Abaqus Verification Manual
- “**VUMAT**: rotating cylinder,” Section 4.1.37 of the Abaqus Verification Manual

### Overview

---

Utility routines **STDB\_ABQERR** and **XPLB\_ABQERR** can be called from any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, to issue an informational, a warning, or an error message to the message (**.msg**) file in Abaqus/Standard or the status (**.sta**) file in Abaqus/Explicit.

### Interface

---

```
DIMENSION INTV(*),REALV(*)
CHARACTER*8 CHARV(*)
...
CALL STDB_ABQERR(LOP,STRING,INTV,REALV,CHARV)
or
CALL XPLB_ABQERR(LOP,STRING,INTV,REALV,CHARV)
...
```

### Variables to be provided to the utility routine

---

#### LOP

Flag for the type of message to be issued.

Set **LOP** = 1 if an informational message is to be issued.

Set **LOP** = -1 if a warning message is to be issued.

Set **LOP** = -2 if an error message is to be issued and the analysis is to be continued.

Set **LOP** = -3 if an error message is to be issued and the analysis is to be stopped immediately.

#### STRING

A string of at most 500 characters long between single quotes containing the message to be issued. If the string needs to be written on more than one line, several one line long strings (between single quotes) should be concatenated using the double forward slash (//) operator.

Integer, real, and character variables can be referenced inside the message using the %I, %R, and %S inserts, respectively. The integer, real, or character variables are passed into the utility routine via

## PRINTING MESSAGES

the **INTV**, **REALV**, and **CHARV** variables, respectively. The variables are then output in the order they are stored in these arrays.

### **INTV**

Array of integer variables to be output. The first %I in **STRING** will output **INTV(1)**, the second **INTV(2)**, and so on.

### **REALV**

Array of real variables to be output. The first %R in **STRING** will output **REALV(1)**, the second **REALV(2)**, and so on.

### **CHARV**

Array of at most 8 character long variables to be output. The first %S in **STRING** will output **CHARV(1)**, the second **CHARV(2)**, and so on.

## 2.1.15 TERMINATING AN ANALYSIS

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “User subroutines: overview,” Section 15.1.1 of the Abaqus Analysis User’s Manual
- “**UMAT** and **UHYPER**,” Section 4.1.21 of the Abaqus Verification Manual
- “**UWAVE** and **UEXTERNALDB**,” Section 4.1.27 of the Abaqus Verification Manual
- “**VUMAT**: rotating cylinder,” Section 4.1.37 of the Abaqus Verification Manual

### Overview

---

Utility routines **XIT** and **XPLB\_EXIT** can be called from within any Abaqus/Standard or Abaqus/Explicit user subroutine, respectively, (except **UEXTERNALDB**) to terminate an analysis. **XIT** or **XPLB\_EXIT** should be used instead of **STOP** to ensure that all files associated with the analysis are closed properly.

### Interface

---

```
CALL XIT
or
CALL XPLB_EXIT
```



## 2.1.16 OBTAINING SENSOR INFORMATION

**Products:** Abaqus/Standard Abaqus/Explicit

### References

---

- “UAMP,” Section 1.1.19
- “VUAMP,” Section 1.2.7
- “Crank mechanism,” Section 4.1.2 of the Abaqus Example Problems Manual

### Overview

---

Utility routines **IGETSENSORID** and **GETSENSORVALUE** can be called only from Abaqus/Standard user subroutine **UAMP**. Utility routines **IVGETSENSORID** and **VGETSENSORVALUE** can be called only from Abaqus/Explicit user subroutine **VUAMP**.

Given the user-defined name for a sensor, these utility routines can be used to obtain the sensor ID or the sensor value using a computationally efficient searching technique.

### Interface

---

```

character*80 mySensorName
...
iMySensorID = IGETSENSORID(mySensorName, jSensorLookUpTable )
iMySensorID = IVGETSENSORID(mySensorName, jSensorLookUpTable )
dMySensorValue = sensorValues(iMySensorID)
...
dMySensorValue = GETSENSORVALUE(mySensorName,
C           jSensorLookUpTable, sensorValues )
C           dMySensorValue = VGETSENSORVALUE(mySensorName,
C           jSensorLookUpTable, sensorValues )
...

```

### Variables to be provided to the utility routine

---

#### **mySensorName**

User-defined character string, uppercase, left justified.

#### **jSensorLookUpTable**

Pointer to an object containing a binary tree look up table for sensors. The calling user subroutine provides this variable.

## OBTAINING SENSOR INFORMATION

### **sensorValues**

Array containing the latest sensor values for all sensors in the model.

### **Variables returned from the utility routine**

---

#### **iMySensorID**

Index in the **sensorValues** array for this sensor name.

#### **dMySensorValue**

Sensor value for this sensor name.

## 2.1.17 ACCESSING Abaqus MATERIALS

**Product:** Abaqus/Standard

### References

---

- “UELMAT,” Section 1.1.25
- “UELMAT,” Section 4.1.15 of the Abaqus Verification Manual

### Overview

---

Utility routine **MATERIAL\_LIB\_MECH** can be called only from Abaqus/Standard user subroutine **UELMAT**.

The routine returns the stress and the material Jacobian at the element material point.

### Interface

---

```
dimension stress(*),ddsdde(ntens,*),stran(*),dstran(*),
* defGrad(3,3),predef(nprefdf),dpredef(nprefdf),coords(3)
...
call material_lib_mech(materiallib,stress,ddsdde,stran,dstran,
* npt,dvdv0,dvmat,dfgrd,predef,dpredef,nprefdf,celent,coords)
...
```

### Variables to be provided to the utility routine

---

#### **materiallib**

Variable containing information about the Abaqus material. This variable is passed into user subroutine **UELMAT**.

#### **stran**

Strain at the beginning of the increment.

#### **dstran**

Strain increment.

#### **npt**

Integration point number.

#### **dvdv0**

Ratio of the current volume to the reference volume at the integration point.

### **dvmat**

Volume at the integration point.

### **dfgrd**

Array containing the deformation gradient at the end of the increment.

### **predef**

Array of interpolated values of predefined field variables at the integration point at the start of the increment.

### **dpredef**

Array of increments of predefined field variables.

### **npredef**

Number of predefined field variables, including temperature.

### **celent**

Characteristic element length.

### **coords**

An array containing the coordinates of this point. These are the current coordinates if geometric nonlinearities are accounted for during the step (see “Procedures: overview,” Section 6.1.1 of the Abaqus Analysis User’s Manual); otherwise, the array contains the original coordinates of the point.

## **Variables returned from the utility routine**

---

### **stress**

Stress tensor at the end of the increment.

### **ddsdde**

Jacobian matrix of the constitutive model,  $\partial\Delta\sigma/\partial\Delta\varepsilon$ , where  $\Delta\sigma$  are the stress increments and  $\Delta\varepsilon$  are the strain increments. **ddsdde(i,j)** defines the change in the *i*th stress component at the end of the time increment caused by an infinitesimal perturbation of the *j*th component of the strain increment array.

## 2.1.18 ACCESSING Abaqus THERMAL MATERIALS

**Product:** Abaqus/Standard

### References

---

- “UELMAT,” Section 1.1.25
- “UELMAT,” Section 4.1.15 of the Abaqus Verification Manual

### Overview

---

Utility routine **MATERIAL\_LIB\_HT** can be called only from Abaqus/Standard user subroutine **UELMAT**.

The routine returns heat fluxes, internal energy time derivative, volumetric heat generation rate, and their derivatives at the element material point.

### Interface

---

```

dimension predef(npredef),dpredef(npredef),dtemdx(*),
*           rhodUdg(*),flux(*),dfdt(*),dfdg(ndim,*),drpldt(*),
*           coords(3)
*
*
call material_lib_ht(materiallib,rhoUdot,rhodUdt,rhodUdg,
*           flux,dfdt,dfdg,rpl,drpldt,npt,dvmat,predef,
*           dpredef,npredf,temp,dtemp,dtemdx,celent,coords)
*
*

```

### Variables to be provided to the utility routine

---

#### **materiallib**

Variable containing information about the Abaqus material. This variable is passed into user subroutine **UELMAT**.

#### **npt**

Integration point number.

#### **dvmat**

Volume at the integration point.

#### **predef**

Array of interpolated values of predefined field variables at the integration point at the start of the increment.

## ACCESSING Abaqus THERMAL MATERIALS

### **dpredef**

Array of increments of predefined field variables.

### **npredf**

Number of predefined field variables, including temperature.

### **temp**

Temperature at the integration point at the start of the increment,  $\theta$ .

### **dtemp**

Increment of temperature.

### **dtemdx**

Spatial gradients of temperature,  $\partial\theta/\partial\mathbf{x}$ , at the end of the increment.

### **celent**

Characteristic element length.

### **coords**

The array containing the original coordinates of this point.

## Variables returned from the utility routine

---

### **rhoudot**

Time derivative of the internal thermal energy per unit mass,  $U$ , multiplied by density at the end of increment.

### **rhodUdt**

Variation of internal thermal energy per unit mass with respect to temperature multiplied by density evaluated at the end of the increment.

### **rhodUdg**

Variation of internal thermal energy per unit mass with respect to the spatial gradients of temperature,  $\partial U/\partial(\partial\theta/\partial\mathbf{x})$ , multiplied by density at the end of the increment.

### **flux**

Heat flux vector,  $\mathbf{f}$ , at the end of the increment.

### **dfdt**

Variation of the heat flux vector with respect to temperature,  $\partial\mathbf{f}/\partial\theta$ , evaluated at the end of the increment.

### **dfdg**

Variation of the heat flux vector with respect to the spatial gradients of temperature,  $\partial\mathbf{f}/\partial(\partial\theta/\partial\mathbf{x})$ , at the end of the increment

**rp1**

Volumetric heat generation per unit time at the end of the increment.

**drpldt**

Variation of **rp1** with respect to temperature.



## **Appendix A: Index**

- “User subroutines index,” Section A.1
- “User subroutine functions listing,” Section A.2



## A.1 User subroutines index

The following tables categorize each user subroutine according to its primary function. The topics are listed alphabetically.

**Table A-1** Abaqus/Standard user subroutines.

Function	Related user subroutines
Amplitudes, User-defined	<b>UAMP</b>
Boundary Conditions	<b>DISP</b>
Constraints	<b>MPC</b>
Contact Behavior	<b>FRIC, FRIC_COEF, GAPCON, GAPELECTR, UINTER</b>
Contact Surfaces	<b>RSURFU</b>
Element Output	<b>UVARM</b>
Elements, User-defined	<b>UEL, UELMAT</b>
Fields, Predefined	<b>UFIELD, UMASFL, UPRESS, USDFLD, UTEMP</b>
Initial Conditions	<b>HARDINI, SDVINI, SIGINI, UPOREP, VOIDRI</b>
Interfacing with External Resources	<b>UEXTERNALDB, URDFIL</b>
Loads, Distributed	<b>DLOAD, UTRACLOAD</b>
Loads, Thermal	<b>FILM, HETVAL</b>
Material Properties	<b>CREEP, UANISOHYPER_INV, UANISOHYPER_STRAIN, UDMGINI, UEXPAN, UFLUID, UFLUIDLEAKOFF, UHARD, UHYPEL, UHYPER, UMULLINS, UTRS</b>
Materials, User-defined	<b>UMAT, UMATHT</b>
Motion, Prescribed	<b>UMESHMOTION, UMOTION</b>
Orientation	<b>ORIENT</b>
Pore Fluid Flow	<b>DFLOW, DFLUX, FLOW</b>
Random Response	<b>UCORR, UPSD</b>
Shell Section Behavior	<b>UGENS</b>
Wave Kinematics	<b>UWAVE</b>

**Table A–2** Abaqus/Explicit user subroutines.

<b>Function</b>	<b>Related user subroutines</b>
Amplitudes, User-defined	<b>VUAMP</b>
Boundary Conditions	<b>VDISP</b>
Contact Behavior	<b>VFRIC, VFRIC_COEF, VFRICTION, VUINTER, VUINTERACTION</b>
Elements, User-defined	<b>VUEL</b>
Fields, Predefined	<b>VUFIELD, VUSDFLD</b>
Fluid Exchange, User-defined	<b>VUFLUIDEXCH, VUFLUIDEXCHEFFAREA</b>
Loads, Distributed	<b>VDLOAD</b>
Material Properties	<b>VFABRIC, VUANISOHYPER_INV, VUANISOHYPER_STRAIN, VUHARD, VUTRS, VUVISCOSITY</b>
Materials, User-defined	<b>VUMAT</b>

## A.2 User subroutine functions listing

The following tables describe the function of each available user subroutine.

### Abaqus/Standard User Subroutines

Name	Function
<b>CREEP</b>	User subroutine to define time-dependent, viscoplastic behavior (creep and swelling).
<b>DFLOW</b>	User subroutine to define nonuniform pore fluid velocity in a consolidation analysis.
<b>DFLUX</b>	User subroutine to define nonuniform distributed flux in a heat transfer or mass diffusion analysis.
<b>DISP</b>	User subroutine to specify prescribed boundary conditions.
<b>DLOAD</b>	User subroutine to specify nonuniform distributed loads.
<b>FILM</b>	User subroutine to define nonuniform film coefficient and associated sink temperatures for heat transfer analysis.
<b>FLOW</b>	User subroutine to define nonuniform seepage coefficient and associated sink pore pressure for consolidation analysis.
<b>FRIC</b>	User subroutine to define frictional behavior for contact surfaces.
<b>FRIC_COEF</b>	User subroutine to define the frictional coefficient for contact surfaces.
<b>GAPCON</b>	User subroutine to define conductance between contact surfaces or nodes in a fully coupled temperature-displacement analysis or pure heat transfer analysis.
<b>GAPELECTR</b>	User subroutine to define electrical conductance between surfaces in a coupled thermal-electrical analysis.
<b>HARDINI</b>	User subroutine to define initial equivalent plastic strain and initial backstress tensor.
<b>HETVAL</b>	User subroutine to provide internal heat generation in heat transfer analysis.
<b>MPC</b>	User subroutine to define multi-point constraints.
<b>ORIENT</b>	User subroutine to provide an orientation for defining local material directions or local directions for kinematic coupling constraints or local rigid body directions for inertia relief.
<b>RSURFU</b>	User subroutine to define a rigid surface.
<b>SDVINI</b>	User subroutine to define initial solution-dependent state variable fields.
<b>SIGINI</b>	User subroutine to define an initial stress field.
<b>UAMP</b>	User subroutine to specify amplitudes.
<b>UANISOHYPER_INV</b>	User subroutine to define anisotropic hyperelastic material behavior using the invariant formulation.

## APPENDIX A: INDEX

<b>UANISOHYPER_STRAIN</b>	User subroutine to define anisotropic hyperelastic material behavior based on Green strain.
<b>UCORR</b>	User subroutine to define cross-correlation properties for random response loading.
<b>UDMGINI</b>	User subroutine to define the damage initiation criterion.
<b>UEL</b>	User subroutine to define an element.
<b>UELMAT</b>	User subroutine to define an element with access to Abaqus materials.
<b>UEXPAN</b>	User subroutine to define incremental thermal strains.
<b>UEXTERNALDB</b>	User subroutine to manage user-defined external databases and calculate model-independent history information.
<b>UFIELD</b>	User subroutine to specify predefined field variables.
<b>UFLUID</b>	User subroutine to define fluid density and fluid compliance for hydrostatic fluid elements.
<b>UFLUIDLEAKOFF</b>	User subroutine to define the fluid leak-off coefficients for pore pressure cohesive elements.
<b>UGENS</b>	User subroutine to define the mechanical behavior of a shell section.
<b>UHARD</b>	User subroutine to define the yield surface size and hardening parameters for isotropic plasticity or combined hardening models.
<b>UHYPEL</b>	User subroutine to define a hypoelastic stress-strain relation.
<b>UHYPER</b>	User subroutine to define a hyperelastic material.
<b>UINTER</b>	User subroutine to define surface interaction behavior for contact surfaces.
<b>UMASFL</b>	User subroutine to specify prescribed mass flow rate conditions for a convection/diffusion heat transfer analysis.
<b>UMAT</b>	User subroutine to define a material's mechanical behavior.
<b>UMATHT</b>	User subroutine to define a material's thermal behavior.
<b>UMESHMOTION</b>	User subroutine to specify mesh motion constraints during adaptive meshing.
<b>UMOTION</b>	User subroutine to specify motions during cavity radiation heat transfer analysis or steady-state transport analysis.
<b>UMULLINS</b>	User subroutine to define damage variable for the Mullins effect material model.
<b>UPOREP</b>	User subroutine to define initial fluid pore pressure.
<b>UPRESS</b>	User subroutine to specify prescribed equivalent pressure stress conditions.
<b>UPSD</b>	User subroutine to define the frequency dependence for random response loading.
<b>URDFIL</b>	User subroutine to read the results file.
<b>USDFLD</b>	User subroutine to redefine field variables at a material point.
<b>UTEMP</b>	User subroutine to specify prescribed temperatures.
<b>UTRACLOAD</b>	User subroutine to specify nonuniform traction loads.

<b>UTRS</b>	User subroutine to define a reduced time shift function for a viscoelastic material.
<b>UVARM</b>	User subroutine to generate element output.
<b>UWAVE</b>	User subroutine to define wave kinematics for an Abaqus/Aqua analysis.
<b>VOIDRI</b>	User subroutine to define initial void ratios.

**Abaqus/Explicit User Subroutines**

<b>Name</b>	<b>Function</b>
<b>VDISP</b>	User subroutine to specify prescribed boundary conditions.
<b>VDLOAD</b>	User subroutine to specify nonuniform distributed loads.
<b>VFABRIC</b>	User subroutine to define fabric material behavior.
<b>VFRIC</b>	User subroutine to define frictional behavior for contact surfaces.
<b>VFRIC_COEF</b>	User subroutine to define the frictional coefficient for contact surfaces.
<b>VFRICITION</b>	User subroutine to define frictional behavior for contact surfaces.
<b>VUAMP</b>	User subroutine to specify amplitudes.
<b>VUANISOHYPER_INV</b>	User subroutine to define anisotropic hyperelastic material behavior using the invariant formulation.
<b>VUANISOHYPER_STRAIN</b>	User subroutine to define anisotropic hyperelastic material behavior based on Green strain.
<b>VUEL</b>	User subroutine to define an element.
<b>VUFIELD</b>	User subroutine to specify predefined field variables.
<b>VUFLUIDEXCH</b>	User subroutine to define the mass flow rate/heat energy flow rate for fluid exchange.
<b>VUFLUIDEXCHEFFAREA</b>	User subroutine to define the effective area for fluid exchange.
<b>VUHARD</b>	User subroutine to define the yield surface size and hardening parameters for isotropic plasticity or combined hardening models.
<b>VUINTER</b>	User subroutine to define the interaction between contact surfaces.
<b>VUINTERACTION</b>	User subroutine to define the contact interaction between surfaces with the general contact algorithm.
<b>VUMAT</b>	User subroutine to define material behavior.
<b>VUSDFLD</b>	User subroutine to redefine field variables at a material point.
<b>VUTRS</b>	User subroutine to define a reduced time shift function for a viscoelastic material.
<b>VUVISCOSITY</b>	User subroutine to define the shear viscosity for equation of state models.

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