Scientific Programming With Modern Fortran

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High Performance Computing I, 2014

Part I

Fortran Basic Operations

History of FORTRAN

One of the earliest of the high-level programming languages, FORTRAN (short for **For**mula **Tran**slation) was developed by John Backus at IBM in the 1950s (circa 1953 for the 704, first released in 1957):

- FORTRAN 66, ANSI standard (X3.9-1966) based on FORTRAN IV
- FORTRAN 77, X3.9-1978, improved I/O
- Fortran 90, ISO/IEC 1539:1991(E), FORTRAN becomes Fortran
- Fortran 95, ISO/IEC 1539-1:1997, minor revisions
- Fortran 2003, ISO/IEC 1539:2004, command line arguments, more intrinsics, IEEE exception handling, C interoperability
- Fortran 2008, revision to Fortran 2003, to include BIT type and Co-array parallel processing (starting to become better supported)

Why Is This Language Still Used?

So why is Fortran still in use?

- Efficiency has always been a priority, in the sense that compilers should produce efficient code. Fortran data are generally not allowed to be aliased (e.g. pointers) making life easier on the compiler
- Designed from the outset as the tool for numerically intensive programming in science and engineering
- Backward compatibility most modern compilers can still compile old (as in decades!) code

Come On, Fortran is 50 Years Old!

Indeed, Fortran celebrated its 50th birthday in 2007. How many other high-level languages can you say that about? Think of it this way - it is arguably the most efficient, compact, portable high-level language available. And you will likely still be able to use programs written today until you retire (unmodified, at that) ...

Some Fortran Compilers

Fortran compilers are easy to find, some are freely available (including compiler source code):

- gfortran, part of the GNU toolchain (gcc/g++), supplanted g77 as of version 4.0
- q95, another freebie
- open64, open sourced by SGI (derived from their MIPSPro compiler suite), further developed by Intel and the Chinese Academy of Sciences (Gnu Public License), intended mainly for compiler research (includes C/C++) on the IA64 platform
- ifort, Intel's commercial compiler
- pgf95, PGI's commercial compiler
- Other commercial compilers: IBM, Qlogic/Pathscale, Lahey/Fujitsu, Sun, NAGWare, ...

The Rest of This Talk

In this presentation I will focus almost entirely on Fortran >=90/95 (a.k.a *Modern Fortran*) features, with some side notes on FORTRAN 77 (mostly for contrast). This talk is not intended to be exhaustive in its coverage of Fortran syntax, but should be enough to yield a "rough and ready" knowledge for HPC.

Source Code Format

While FORTRAN 77 used by default a **fixed** format for its source code, the default with modern Fortran is now a **free** format, in which lines can be up to 132 columns long, with a maximum of 39 **continuation** lines, indicated by the & character.

Source Format Summary

```
132 characters per line
! initiates comment
& continuation character
; statement separator (multiple per line)
```

```
print*,''You need two continuation &
    &characters when splitting a token &
    & or string across lines''
```

Statement Labels

Statement labels are still in fashion, and consist of 1-5 digits (leading zeros are neglected), e.g.

```
1 101 WRITE(6,*) 'Enter runid:(<=9chars) '
READ(*,2) runid
2 format (a9)
```

Most commonly used for FORMAT statements.

Names

In Fortran, names must:

- Start with a letter (a-z)
- Contain only letters, digits, and underscore
- Must not be longer than 31 characters

More Resources

Good reference material on all things Fortran:

- M. Metcalf and J. Reid, Fortran 90/95 Explained, 2nd Ed., (Oxford, Great Britain, 2000)
 Fortran 95/2003 Explained, 3rd Ed., (Oxford, Great Britain, 2004), with M. Cohen
 Modern Fortran Explained, (Oxford, Great Britain, 2011), with M. Cohen
- Metcalf's online tutorial:

```
http://wwwasdoc.web.cern.ch/wwwasdoc/WWW/f90/f90.html
```

• Alan Miller's Fortran Resources:

```
http://jblevins.org/mirror/amiller/
```

Implicit Types

Unfortunately FORTRAN historically allowed **implicit types**, which by default were:

FORTRAN 77

implicit real (a-h,o-z) implicit integer (i-n)

Thanks to backward compatibility, this feature is still present:

Fortran

implicit type (letter-list) [,type (letter-list) ...]

Implicit None

I would recommend starting every declaration block with the following:

Fortran

implicit none

which will turn all implicit typing off, and save a lot of potential errors (misspelled variable names just being one of them). Use implicit types at your own risk!

Basic Declarations

Declarative Syntax

<type> [,<attribute-list>] :: <variable-list> & [=<value>]

The attribute is usually one of PARAMETER, DIMENSION, or POINTER.

Fortran Intrinsic Types

We have five different intrinsic types, integer, real, logical, complex, and character. The syntax looks like:

```
1
integer :: i
real :: x
3 complex :: z
4 logical :: beauty
character :: c
```

These are all of default **kind** ... (Note that integers can also be represented in binary (base 2), octal (base 8), and hex (base 16))

Kind

There is an intrinsic function KIND that returns an integer and can be used to query (or set) the "kind type" of a variable

Fortran

KIND(x) returns default integer whose value is the kind type parameter value of x

Example: Declarative Use of KIND

These are two of the very handy "transformational" functions:

SELECTED_REAL_KIND/SELECTED_INT_KIND

SELECTED_REAL_KIND([p][,r])

- p:: integer, decimal precision (intrinsic PRECISION)
- r :: integer, decimal exponent range (RANGE)
 (at least one of p or r must be present)

SELECTED_INT_KIND(r)

r :: integer, decimal exponent range

the return values of both functions are default integers whose values can be used in the KIND intrinsic function. -1 is returned if the desired precision is unavailable (-2 for the range in SELECTED_REAL_KIND, -3 if both).

Best illustrated by example - this is a chunk of code that use in all of my Fortran codes:

```
integer, parameter :: si=KIND(1), sp=KIND(1.0), sc=KIND((1.0,1.0)), &

di=SELECTED_INT_KIND(2*RANGE(1_si)), &

dp=SELECTED_REAL_KIND(2*PRECISION(1.0_sp)), &

dc=SELECTED_REAL_KIND(2*PRECISION(1.0_sc))
```

Note the use of underscores in literal constants to indicate kind type, which is quite generally applicable:

PARAMETER Attribute

Instead of the old FORTRAN 77 PARAMETER statement, one can use the parameter attribute:

```
REAL, PARAMETER :: E = 2.71828, PI = 3.141592
```

Intrinsic Type Arrays

Arrays can be indicated by the old-type DIMENSION attribute, or simply in the declaration itself:

are all arrays of rank 1, shape 3.

Note that arrays can be sectioned:

```
1 a(i:j) ! rank 1 array, size j-i+1
2 b(k,1:n) ! rank 1 array, size n
3 c(1:m,1:n,k) ! rank 2 array, extent m by n
```

and vector subscripts can be used:

```
1 x(ivector) ! ivector is an integer array
```

which makes for a very flexible (essentially arbitrary) indexing scheme.

Initializing Arrays

There is some special syntax for initialization arrays:

```
1 a(1:4) = 0.0
2 a(1:4) = (/1.1, 1.2, 1.3, 1.4/)
3 a(1:4) = (/ i, i=1,7,2 /) ! implied do loop with stride 2
```

Note that the RESHAPE function can also be used to initialize an array of rank greater than 1.

Character Arrays

Better known as **strings**, Fortran is not the best for string handling (it is intended for number crunching, after all), but modern Fortran is considerably improved in this regard:

Substrings can be handled in several ways:

```
character(len-1200) :: line
character(len=12) :: words,term*8 ! term has length 8, alternate length spec
line(:i) ! same as line(1:i)
line(i:) ! same as line(i:120)
line(:) ! same as line(1:120)
```

and we will come back later to some of the intrinsic string handling functions and subroutines.

Derived Types

It is often advantageous to define your own data types:

```
1
2
    TYPE Coords_3D
2    real :: r,theta,phi ! spherical coordinates
3
4
5
TYPE (Coords_3D)
TYPE (Coords_3D) :: point1, point2
```

You can reference the internal components of a structure:

```
1 x1 = point1%r * SIN(point1%theta) * COS(point1%phi)
2 y1 = point1%r * SIN(point1%theta) * SIN(point1%phi)
3 z1 = point1%r * COS(point1%theta) *
4 x2 = point2%r * SIN(point2%theta) * COS(point2%phi)
5 y2 = point2%r * SIN(point2%theta) * SIN(point2%phi)
6 z2 = point2%r * COS(point2%theta)
```

N.B. Derived type components can not be ALLOCATABLE (but they can be POINTERS).

Assigning Derived Types

Derived types can be assigned either by component or using a **constructor**:

```
point1%r = 1.0
point1%theta = 0.0
point1%phi = 0.0
point1 = Coords_3D(1.0,0.0,0.0)
point2 = point1
```

Note that two variables of the same type can be handled very simply. Derived types should always be placed in a **MODULE**.

Pointers

Fortran finally has the capability (often best left unused) of using pointers. Along with ALLOCATABLE and automatic arrays, pointers make up the 3 types of dynamic data in Fortran 90/95. Pointers can be members of derived types, most useful for implementation of linked lists:

We will talk a bit more about pointers later when dealing with other aspects of dynamic memory.

DATA Statement

The DATA statement can be used to initialize values:

```
1 DATA object-list /value-list/ [[,] object-list /value-list/] ...
```

Values initialized in DATA statements automatically have the SAVE attribute.

Scalar Expressions

Scalar expressions use operators **, *, /, +, -. The following table summarizes the result KIND for all but exponentiation (**)

а	b	used(a)	used(b)	result
П		а	b	
I	R	real(a,kind(b))	b	R
I	С	cmplx(a, 0, kind(b))	b	С
R	-1	a	real(b,kind(a))	R
R	R	a	b	R
R	С	cmplx(a, 0, kind(b))	b	С
С	-1	a	cmplx(b, 0, kind(a))	С
С	R	а	cmplx(b, 0, kind(a))	С
С	С	a	b	С

(I = integer, R = real, C = complex)

and for exponentiation (**):

а	b	used(a)	used(b)	result
I		а	b	
I	R	real(a,kind(b))	b	R
1	С	cmplx(a, 0, kind(b))	b	С
R	-1	a	b	R
R	R	a	b	R
R	С	cmplx(a, 0, kind(b))	b	С
С	-1	a	b	С
С	R	a	cmplx(b, 0, kind(a))	С
С	С	a	b	С

(I = integer, R = real, C = complex)

Relational Operators

Fortran supports the old-style FORTRAN 77 relational operators as well as more modern syntax (which most compilers were supporting by extension anyway):

77	Modern
.lt.	<
.le.	<=
.eq.	==
.ne.	/=
.gt.	>
.ge.	>=

Logical Operators

The logical operators are simply:

```
.not. logical negation
```

.and. logical intersection

.or. logical union

.eqv. logical equivalence

.neqv. logical non-equivalence

Character Operations

Substrings are easy using the same syntax as for arrays:

where we also made use of the "//" concatenation operator.

Array Assignment

Arrays can be assigned using expressions that involve arrays of the same shape, or by scalar (in which case the scalar is applied to all elements of the array), e.g.

```
1 real :: a(20,20),b(10)
2 a(1,11:20) = b
a = a + 1.0
```

This flexible syntax makes for much more compact code.

IF Construct

```
1 [name:] if (expr) then
2 block
3 [else if (expr) then [name]
4 block] ...
5 [else [name]
6 block]
7 end if [name]
```

Note that the optional name is only for program clarity (especially for nested loops), and needs to be unique. Also note the single statement variation:

```
1 if (expr) single-statement
```

DO Loops

Old style loop syntax:

```
1 [name:] do var=expr1,expr2[,expr3]
2 block
end do [name]
```

where the loop is executed MAX(0,(expr2-expr1+expr3)/expr3) times, and var is a named scalar integer variable, and expr1, expr2, and expr3 are scalar integer expressions (expr3 must be nonzero if used).

Infinite DO Loop

One important variation on the DO construct is the endless do loop, which has a simple means to exit:

```
1 2 i = i+1 3 if (i >= enough) exit end do
```

Note that the <code>exit</code> statment is also handy for early termination in bounded loops. The <code>cycle</code> statement is similar, but just drops execution to the next iteration.

DO-WHILE Loop?

There is a DO-WHILE construct, but with the exit option for early loop termination, it is not really worth talking about ...

```
1 2 ... (a /= b) ... END DO
```

is exactly the same as

The GOTO Statement

Ah, the bane of FORTRAN 77 programmers everywhere - the infamous GOTO statement. Still supported in Fortran 90/95, be wary of using the GOTO. If you have to use it, make it as clear as you possibly can.

```
1 101 WRITE(6,*) 'Enter runid:(<=9chars) '
READ(*,'(a9)') runid
3 i=INDEX(runid,' ')-1
4 INQUIRE(file=runid(1:i)//".in",exist=ifex)
5 if(.not.ifex) goto 101
OPEN(unit=iunit,file=runid(1:i)//'.in',status='old', &
form='formatted')
```

Using Named and Nested Loops

Here is an example of when using named loops comes in handy:

```
1 outa: DO
2 inna: DO
3 ...
4 IF (a.GT.b) EXIT outa ! jump to line 10
5 IF (a.EQ.b) CYCLE outa ! jump to line 1
6 IF (c.GT.d) EXIT inna ! jump to line 9
7 IF (c.EQ.a) CYCLE inna ! jump to line 2
8 END DO inna
END DO outa
```

The CASE Construct

Somewhat like the C switch statement ...

```
The CASE construct can be used as an efficient substitute for a more elaborate IF ... THEN ... ELSEIF ... ELSE ... END IF construct.
```

An example of using CASE when parsing an input file:

Part II

Fortran Essentials

Modules

Modules provide a way to package together commonly used code (similar to the old common blocks in FORTRAN 77) and have distinct advantages:

- Can be used to hide internal data and routines through PRIVATE and PUBLIC declarations
- Can contain common subroutines and functions with explicit interfaces which can be changed without affecting calling code
- Modules can (and often should) be compiled separately, before the program units that use them

Module Syntax

```
module module-name
[specification statements]
[contains
module-subprograms]
end [module [module-name]]
```

and a very simple example:

7

Using Modules

Modules encapsulate code that can be made accessible to other program units through the USE statement:

```
1 MODULE TBatoms
2 USE TBconst ...
```

Modules are free to load other modules, but not themselves.

Module Data Visibility

You can allow or prevent access to the internal workings of a module using the PRIVATE and PUBLIC attributes:

```
PRIVATE :: pos, store, stack size
                                            ! hidden
        PUBLIC
                :: pop, push
                                            ! not hidden
or
```

```
PUBLIC
                       ! set default visibility
INTEGER, PRIVATE, SAVE :: store(stack size), pos
INTEGER.
        PRIVATE, PARAMETER :: stack size = 100
```

Renaming/USE ONLY

You can rename a module entity in a local context:

```
1 USE TBconst, local_dp => dp    ! dp becomes local_dp in current scope
```

or you can restrict access by

```
1 USE TBconst, ONLY:dp,dc ! only load dp and dc from module TBconst
```

Module Summary

```
MODULE modname
 2
      ... type defs
      ... Global data
 5
      CONTAINS
       SUBROUTINE sub 1
 7
       CONTAINS
         SUBROUTINE internal 1
10
11
         END SUBROUTINE internal 1
12
         SUBROUTINE internal 2
13
14
         END SUBROUTINE internal 2
15
       END SUBROUTINE sub 1
16
17
       FUNCTION fun_1
18
19
       CONTAINS
20
21
       END FUNCTION fun 1
22
     END MODULE modname
```

Subroutine Syntax

The syntax for a subroutine call is given by

SUBROUTINE

[recursive] subroutine subroutine-name[([dummy arguments])]

Argument Intent

You can provide information to the compiler (always a good idea!) about the dummy arguments to a routine by:

```
1 REAL, INTENT(IN) :: arg1 ! passing in value
2 REAL, INTENT(OUT) :: arg2 ! returning value only
REAL, INTENT(INOUT) :: arg3 ! both apply
```

You should always make use of the INTENT attribute - it allows the compiler to do extensive error checking and optimization (remember, the more that your compilers knows about your code, the better it will be able to perform).

SAVE Attribute

SUBROUTINE sub 1 (arg1, arg2)

The SAVE attribute can be applied to a specified entity, or all of the local entities in a procedure:

```
2    integer, save :: number_calls = 0
    ...
    number_calls = number_calls +1

1    SUBROUTINE sub_1(arg1, arg2)
    ...
2    SAVE
```

SAVE acts to preserve values between calls to the subroutine/function.

Function Call Syntax

The syntax for a function call is given by

FUNCTION

type [recursive] function *function-name*[([dummy arguments])] & [result(result-name)]

Explicit Interfaces

External subprograms have an implicit interface by default (even if one uses the <code>external</code> statement to indicate that a subunit is outside the current code, the arguments and their types remain unspecified), and an <code>INTERFACE</code> block is necessary to specify an explicit interface of an external subprogram; as mentioned above, this allows type-checking of actual and formal arguments in a reference to a subprogram

Explicit Interface Example

SUBROUTINE resid(m, n, x, fvec. iflag)

```
3
            USE TBatoms
            USE TBconst
            USE TBfitdata
            USE TBflags
            USE TBmat
            USE TBopt
            USE TBparams
10
11
            implicit none
12
            integer, intent(in) :: m, n
13
            integer, intent(in out) :: iflag
14
            real(kind=dp), intent(in) :: x(n)
15
            real(kind=dp), intent(in out) :: fvec(m)
16
          END SUBROUTINE resid
17
       END INTERFACE
```

Note that the syntax of the interface-body is just an exact copy of the subprogram's header, argument specifications, function result, and END statement.

INTERFACE

INTERFACE Properties

- Can not use both EXTERNAL and INTERFACE
- Explicit interfaces required for POINTER or TARGET dummy arguments in a procedure, or pointer-valued function result
- Explicit interfaces also required for OPTIONAL, KEYWORD, and procedural arguments
- Even when not required, explicit interfaces are a good idea (usually placed inside a MODULE)

Optional & Keyword Arguments

Dummy arguments can be optional - using OPTIONAL:

```
1  SUBROUTINE optargs(a,b)
2  REAL, INTENT(IN), OPTIONAL :: a
3  INTEGER, INTENT(IN), OPTIONAL :: b
4  REAL :: ay; INTEGER :: by
5  ay = 1.0; bee = 1 ! defaults
7  IF(PRESENT(a)) ay = a
8  IF(PRESENT(b)) bee = b
9
```

```
CALL optargs()
CALL optargs(1.0,1); CALL optargs(b=1,a=1.0) ! same, using keywords
CALL optargs(1.0); CALL optargs(a=1.0) ! keywords handier still for long lists
CALL optargs(b=1)
```

Note that optional and keyword arguments need explicit interfaces, and should come after positional arguments.

Procedures as Arguments

When using a procedure as an argument, an explicit interface is required (as it is for POINTER, optional, and keyword arguments):

```
REAL FUNCTION minimum(a, b, func)
      returns the minimum value of the function func(x)
    ! in the interval (a,b)
         REAL, INTENT(in) :: a, b
         INTERFACE
             REAL FUNCTION func(x)
                REAL, INTENT(IN) :: x
             END FUNCTION func
         END INTERFACE
10
         REAL f.x
11
12
           = func(x) ! invocation of the user function.
13
14
      END FUNCTION minimum
```

Recursion

Recursion is supported in Fortran >=90/95; we can illustrate it best by example:

```
1 RECURSIVE FUNCTION factorial(n) RESULT (res)
2 INTEGER res, n
3 IF (n.EQ.1) THEN
4 res = 1
5 ELSE
6 res = n*factorial(n-1)
7 END IF
END FUNCTION factorial
```

this would be an example of direct recursion ...

and an example of **indirect** recursion:

```
volume = integrate(func, bounds)
 2
    RECURSIVE FUNCTION integrate(f, bounds)
    ! Integrate f(x) from bounds(1) to bounds(2)
       REAL integrate
       INTERFACE
          FUNCTION f(x)
             REAL f. x
          END FUNCTION f
10
      END INTERFACE
11
       REAL, DIMENSION(2), INTENT(IN) :: bounds
12
13
    END FUNCTION integrate
14
15
    FUNCTION func(x)
16
                      ! module MODfunc contains function f
      USE MODfunc
17
      REAL func, x
18
          x = f = x
19
         func = integrate(f, bounds)
20
    END FUNCTION func
21
```

Generally indirect recursion is of the form 'A calls B calls A ...' - in this case integrate calls func which calls integrate ...

Fortran I/O

Too large a topic to cover in its entirety here - we will focus on the basics required to familiarize you with basic Fortran I/O functionality.

```
OPEN
         ([UNIT=integer, ] FILE=filename, [ERR=label, ] &
2
           [STATUS=status,] [ACCESS=method,] [ACTION=mode,] &
          [RECL=int-expr)
```

- filename is a string
- status is 'OLD'. 'NEW'. 'REPLACE'. 'SCRATCH' or 'UNKNOWN'
- method is 'DIRECT' or 'SEQUENTIAL'
- mode is 'READ', 'WRITE' or 'READWRITE'
- RECL (record length) needs to be specified for DIRECT access files

OPEN Examples

Note that UNIT 1-7 are typically reserved (6, or *, is almost always the standard output, for example), and each file stream needs a unique number.

INQUIRE

A very handy statement for querying the status of a file by unit number or filename:

```
INQUIRE ([UNIT=]unit | FILE=filename, ilist)
```

and there are many possible entries in ilist, of which the most handy are:

- IOSTAT=ios as in the OPEN syntax
- EXIST=log_exist returns a logical on the existence of the file
- OPENED=log_opened returns a logical on whether the file is open

```
!
! Open input file - fail gracefully if not found.
!
INQUIRE(file=runid(1:len_runid)//'.in', exist=exist_in)
if (.not.exist_in) then
write(*,*) '<readin> Unable to open input file: ', &
runid(1:len_runid)//'.in'
stop
endif
OPEN(file=runid(1:len_runid)//'.in', status='old', unit=inunit)
```

Other I/O Statements

CLOSE unattach specified unit number
REWIND place file pointer back to start of file
BACKSPACE place file pointer back one record
ENDFILE force writing end-of-file

READ

where the non-obvious entries are:

- unit is an integer (some lower values are reserved) or * for standard input
- format is a string of FORMAT statement label number
- label is a statement label
- adv-mode is 'YES' or 'NO'
- IOSTAT returns zero for no error

READ Example

```
READ (14,FMT='(3(F10.7,1x))',REC=iexp) a,b,c
READ (14,'(3(F10.7,1x))',REC=iexp) a,b,c ! same as above
READ (*,'(A)',ADVANCE='NO',EOR=12,SIZE=nch) str
```

WRITE

where the entries are as in the READ case.

FORMAT

Fortran does have quite an elaborate formatting system. Here are the highlights.

lw w chars of integer data

Fw.d w chars of real data, d decimal places

Ew.d w chars of real data, d decimal places

Lw w chars of logical data

Aw w chars of CHARACTER data

nX skip n spaces

Note that:

The E descriptor is just the F with scientific notation

FORMAT Example

```
1 WRITE (*, FMT=' (2X,2(I4,1X),''name '', A4,F13.5,1X,E13.5)') & 77778,3,'abcdefghi',14.45,14.5666666

1 **** 3 name abcd 14.45000 0.14567E+02
```

Note that are quite a few other format descriptors, much less commonly used.

Unformatted I/O

Unformatted I/O is simpler (no FORMAT statements) and involves less overhead, less chance of roundoff error), but is inherently non-portable, since it relies on the detailed numerical representation. A file must be either entirely formatted or unformatted, not a mix of the two.

```
1 READ (14) A WRITE (15, IOSTAT=ios, ERR=2001) B
```

Note that unformatted i/o is generally quite a lot faster than formatted - so unless you are concerned with moving your files from one platform to another, you will be much better off using unformatted i/o.

Assumed-shape Arrays

Arrays passed as dummy arguments should generally be what are called **assumed-shape** arrays, meaning that the dimensions are left to the actual (calling arguments):

```
1 SUBROUTINE stubby(a,b)
2 implicit none
3 real,intent(in) :: a(:),b(:,:)
:
```

- Note that the default bounds (1) apply
- Actual arguments can not be vector subscripted or themselves assumed-shape

Automatic Arrays

Local arrays whose extent is determined by dummy arguments are called **automatic objects**. Example:

```
1
2
SUBROUTINE stubby1(b,m,n)
integer, intent(in) :: m,n
3
real, intent(inout) :: b(:,:) ! assumed
REAL :: b1(m,n) ! automatic
5
REAL :: b2(SIZE(b,1),SIZE(b,2)) ! automatic
```

Note that both assumed-shape arrays and automatic objects are likely to be placed on the **stack** in terms of memory storage.

Allocatable Arrays

real, allocatable :: a(:,:)

Finally, dynamic data storage elements for Fortran! An array that is **not** a dummy argument or function can be given the ALLOCATABLE attribute:

```
2
3
4
ALLOCATE (a(ntypes,0:ntypes+2)) ! ntypes is an integer
5
6 ! lots of work
7
8 DEALLOCATE (a)
```

Note that originally ALLOCATABLE arrays could not be part of a derived type (have to use a POINTER to get the same functionality) - that oversight was fixed in Fortran 2003.

ALLOCATE & DEALLOCATE

The syntax for ALLOCATE and DEALLOCATE are given by:

```
1 ALLOCATE( list [,stat=istat] )
DEALLOCATE( list [,stat=istat] )
```

The optional stat= specifier can be used to test the success of the (de)allocation through the scalar integer istat. As usual, zero for success. Leaving out "stat=" should result in a termination if the (de)allocation was unsuccessful.

Pointer Flexibility

Note that you can not associate pointers with just any variable (as in C), instead the variables must be declared using the target attribute:

```
real, target :: x, y(100), z(4,4)
    integer, target :: m, n(10), k(10,10)
 3
 4
    real, pointer :: ptr1, ptr2, ptr_y(:), ptr_z1(:), ptr_z2(:,:)
 5
    ptr1 => x ! simple pointer assignment
    alpha = exp(ptrl) ! pointer shares memory location with x.
                       ! but used like any value
    nullify(ptr1)
                       ! frees up pointer
10
11
    if (associated(ptr1)) then
12
       print*, 'ptrl is associated'
13
       if (associated(ptr1, target=x)) then
14
          print*, 'ptrl is associated with "x"'
15
       endif
16
    endif
```

More Fun With Pointers

array-associated pointers have considerable flexibility:

```
1  ptr_y => y     ! can use ptr_y(i) just as y(i)
2  ptr_y => y(11:20)    ! ptr_y(1) is now y(11) ...
3  ptr_y => z(2,1:4)    ! loads row 2 of z intro ptr_y(:)
4  ptr_z2 => z(2:4,2:4)    ! ptr_z2 is 3x3 submatrix of z
5  ALLOCATE(ptr_z1(16))    ! direct allocation
```

Note that pointers can also be used as components of derived types, making for very flexible data structures.

Pointer Considerations

Some things to think about when using pointers in Fortran:

- Can create complex (and difficult to maintain) code
- Easy to create bugs that only arise at run-time
- Inhibit compiler optimization (difficult to predict data patters and disjoint memory structures)

Elemental Operations

We have already seen **elemental** operations in which conformable operands can be used with intrinsic operators, e.g.

will apply the square root operator individually to all of the elements of a. Not only intrinsics can be elemental, and you can also use the elemental declaration in user-defined functions (Requires Fortran >=95) as well.

Array-valued Functions

Functions can return arrays - just be careful that you ensure that the interface is an explicit one. An example:

```
PROGRAM arrfunc
       implicit none
       integer, parameter :: ndim=36
       integer, dimension (ndim, ndim) :: m1, m2
       m2 = funkv(m1.4)
 7
       CONTAINS
       FUNCTION funky (m, scal)
10
         integer, intent(in) :: ima(:,:)
11
         integer, intent(in) :: scal
12
         integer :: funky(SIZE(m, 1), SIZE(m, 2))
13
         funky(:,:) = m(:,:)*scal
14
       END FUNCTION funky
15
    END PROGRAM arrfunc
```

WHERE Statement

Useful for performing array operations only on certain elements of an array, but preserving the compact syntax:

```
MHERE (logical array-expr)
array-assignments

END WHERE

where (pressure <= 1.0)
pressure = pressure + increment

ELSEWHERE
pressure = pressure + check_pressure

END WHERE</pre>
```

in the example all arrays are of the same shape, and the assignment is said to be *masked* using the comparison (which is done element by element). Fortran >=95 also provides for masks in additional ELSEWHERE statements

FORALL Construct (Fortran 95)

Fortran 95 introduced the concept of a FORALL statement which is basically an array assignment with some explicit indexing:

Implied is that the assignment is trivially data-parallel, i.e. it can be carried out in any order, and therefore can be more efficient than a more traditional loop. Construct form:

where the results are executed in any order, held in temporary storage (to avoid indeterminate results), and then updated in arbitrary order.

Complete FORALL construct syntax:

The body of a FORALL construct can be quite general (containing statements, additional FORALL or WHERE statements/constructs, etc.), but must not branch (e.g. goto) out of the construct. Any included subprograms must be **pure**, in the sense of inducing no undesired side-effects in the sense of inducing an order dependence that would impede parallel processing.

PURE Procedures (Fortran >=95)

Programmer can assert that a function or subroutine is PURE: by adding the PURE keyword to the function/subroutine statement:

- a pure function does not alter its dummy arguments (must be INTENT (IN))
- INTENT of dummy arguments must be declared (IN for functions)
- does not alter variables accessed by host or use association
- contains no local variables with SAVE attribute
- contains no operations on external file
- contains no STOP statements
- any internal procedures must also be pure

all intrinsic functions are pure.

Array Intrinsics

Fortran is designed around the notion of data manipulation, so it is not a great surprise that it has a number of built-in functions for array manipulation, some of which we have already seen (elemental operations, masking).

Array Functions

Set of operations that involve common extractions from arrays:

ALL(MASK [.dim])
ANY(MASK [.dim])
COUNT(MASK [.dim])
MAXLOC(ARRAY [.mask])
MINLOC(ARRAY [.dim[.mask]])
MAXVAL(ARRAY [.dim[.mask]])
MINVAL(ARRAY [.dim[.mask]])
PRODUCT(ARRAY [.dim[.mask]])
SUM (ARRAY [.dim[.mask]])

all relations in mask are true [along dimension dim] if any elements of mask are true [along dimension dim] number of elements of mask that are true location of element with maximum value location of element with minimum value maximum value [of true elements in mask, along dim] minimum value [of true elements in mask, along dim] products [of true elements in mask, along dim] of values sum [of true elements in mask, along dim] of values

Example of Array Extraction Intrinsics

Array Inquiry Functions

Very useful array inquiry functions:

ALLOCATED(ARRAY) LBOUND(ARRAY [,dim]) SHAPE(ARRAY) SIZE(ARRAY [,dim]) UBOUND(ARRAY [,dim]) logical if A has been allocated lower bound for dimension dim of A (integer vector if no dim) returns integer vector of shape of A size of dimension dim of A (else all of A) upper bound for dimension dim of A (integer vector if no dim)

Array Reshaping

RESHAPE is a general intrinsic function which delivers an array of a specific shape:

```
1 RESHAPE(source, shape [,pad][,order])
```

returns an array whose shape is given by the constant rank-1 integer array (nonnegative elements) shape derived from the array source. If order is absent, elements of pad are used to fill out remaining elements in the result (whose size may then exceed that of source. order can be used to pad the result in non-element order.

produces (recall that Fortran is column-ordered):

```
1 A2: 1,1 1,2 = 1.000000 3.000000
A2: 2,1 2,2 = 2.000000 4.000000
```

Vector/Matrix Intrinsics

Fortran 90 has several intrinsics for vector dot products matrix multiplication and transposition:

DOT_PRODUCT(vector_1,vector_2)
MATMUL(matrix_1,matrix_2)
TRANSPOSE(matrix)

dot product of two rank-1 equal length vectors matrix multiplication transposition of any rank-2 array

Intrinsics Categories

There are roughly 75 new intrinsic routines (versus FORTRAN 77), but they roughly fall into 4 categories:

- Elemental procedures
- Inquiry functions
- Transformational functions
- Nonelemental subroutines

I am going to group them a bit differently, and only cover the more common ones. Consult a good reference¹ for a thorough list.

¹Metcalf, Reid, and Cohen *Modern Fortran Explained*, (Oxford, Great Britain, 2011).

Mathematical Intrinsics

Far too many to enumerate here - you can find a handy reference for the full set in other references. Note that almost all support a generic interface supporting available KIND types, and that most are elemental.

```
SQRT, EXP, LOG, LOG10, SIN, COS, TAN, ASIN, ACOS, ATAN, SINH, COSH, TANH
```

Numerical Intrinsics

The list of numerical intrinsics (with syntax and usage):

INT(a[.KIND]) REAL(af.KIND1) CMPLX(x[,y][,KIND]) AINT(a[,KIND]) ANINT(a[.KIND]) NINT(a[,KIND]) ABS(a) MOD(a.p) MODULO(a,p) FLOOR(a[,kind]) CEILING(af.KIND1) SIGN(a.b) DIM(x,y)MAX(a1,a2[,a3,...]) MIN(a1,a2[,a3,...]) AIMAG(z) CONJG(z)

convert to integer, type KIND convert to real, type KIND convert x or (x,y) to complex, type KIND truncate real to lowest whole number, type KIND returns nearest whole number real, type KIND integer (type KIND) value nearest a absolute value of a, same KIND as a remainder of a modulo p, a-int(a/p)*p (has sign of a) a-floor(a/p)*p (has sign of p) greatest integer less than or equal to a, type KIND least integer greater than or equal to a, type KIND absolute value of a times sign of b max(x-y,0.0)maximum of two or more numbers minimum of two or more numbers imaginary part of complex number z, type real, KIND(z) conjugate of complex number z

String Functions

Yes, Fortran does have string handling capability! And in fact, it is much improved. The following table gives a brief synopsis:

ACHAR(I) ADJUSTL(STRING) ADJUSTR(STRING) CHAR(I, kind) IACHAR(C)

ICHAR(C) INDEX(STRING, SUBSTRING, back)

LEN(STRING) LEN TRIM(STRING)

REPEAT(STRING, NCOPIES)

SCAN(STRING, SET, back) TRIM(STRING)

VERIFY(STRING, SET, back)

ASCII character of number I

Adjusts to the left Adjusts to the right

Returns character of number I ASCII number of char C

Number of char C

Starting pos of substring in string

Length of STRING

Length of string without trailing blanks

String concatnation

Position of 1st occurrence of any char in SET in STRING

Returns string without trailing blanks

Position of 1st char in STRING not in SET

The following functions can be used for ASCII lexical string comparisons:

```
LGE (STRING A, STRING B)
LGT (STRING A, STRING B)
LLE (STRING A, STRING B)
LLT (STRING A, STRING B)
```

Note that if the strings are of differing length, the shorter will be padded with blanks for comparative purposes. All return default logical results.

Bitwise Intrinsics

Modern Fortran added support for quite a few bitwise operations:

BIT SIZE(I) number of bits in a word BTEST(I, POS) .true, if POS number of Lis 1 IAND(I, J) logical addition of bit chars in I and J IBCLR(I, POS) puts a zero in the bit in POS IBITS(I, POS, LEN) uses LEN bits of word I beginning at POS, additional bits are set to zero. POS + LEN <= BIT SIZE(I) IBSET(I, POS) puts the bit in position POS to 1 IEOR(I, J) performs logical exclusive OR IOR(I, J) performs logical OR ISHIFT(I, SHIFT) performs logical shift (to the right if the number of steps SHIFT < 0 and to the left if SHIFT > 0). Positions that are vacated are set to zero. performs logical shift a number of steps ISHIFTC(I, SHIFT, size) circularly to the right if SHIFT < 0, circularly to the left if SHIFT > 0. If SIZE

logical complement

is given, it is required that 0 < SIZE <= BIT_SIZE(I)

NOT(I)

Random Numbers

Modern Fortran also has a built-in pseudorandom generator:

```
1 CALL RANDOM_NUMBER(harvest)
CALL RANDOM_SEED([size] | [put] | [get])
```

harvest can be an array, and the range of the random numbers are the interval [0,1). size is intent OUT and returns the size of the integer seed array, which can be input (put) or returned (get).

Real-time Clock

Modern Fortran provides a pair of routines to access the real-time clock:

```
DATE_AND_TIME
```

DATE_AND_TIME([date] [,time] [,zone] [,values])

date character string in form ccyymmdd

time character string in form hhmmss.sss

zone character string in form Shhmm, difference between local and UTC

values integer vector of size at least 8 with year, month, day, difference from UTC in minutes, hour, minutes, seconds, milliseconds

SYSTEM_CLOCK

SYSTEM_CLOCK([count][,count_rate][,count_max])

count processor-dependent value of processor clock
 (-huge(0) if no clock)

count_rate clock counts per second (0 if no clock)

count max maximum for count (0 if no clock)

CPU Time

Fortran >=95 only:

CPU_TIME

CPU_TIME(time)

time real assigned to processor-dependent time in seconds (negative value if no clock)

In my experience the $\mathtt{CPU_TIME}$ intrinsic is not very precise, however, and depends rather strongly on the compiler ...

STOPWATCH

STOPWATCH is not part of standard Fortran, but is a nice little package written by William Mitchell at NIST:

which supports a more full featured set of timing routines (including wall time, cpu time, and system time).

Array Functions

The array intrinsic functions will be discussed in a special section devoted to arrays ...

Part III

Fortran Advanced Operations

Numerical Inquiry Functions

There are a bunch of numerical inquiry functions in Fortran 90/95. First, the integer representation model is given by

$$i = s \sum_{k=0}^{q-1} d_k r^k,$$

where

i integer value

s sign (+,-)

r radix (r > 1)

q number of digits (q > 1)

 d_k k-th digit, $(0 \le d_k < r)$

The floating-point model is given by

$$x = sb^e \sum_{k=1}^p f_k b^{-k},$$

where

```
real value
X
```

$$s$$
 sign $(+,-)$

b base
$$(b > 1)$$

e exponent
$$(q > 1)$$

p number mantissa digits
$$(p > 1)$$

$$f_k$$
 k-th digit, $(0 \le f_k < b)$

Inquiry Functions

The inquiry functions are given in the following table:

digits(x) value of (q, p) for (integer, real)

epsilon(x) b^{1-p}

huge(x) largest value in model

minexponent(x) minimum e maxexponent(x) maximum e

precision(x) decimal precision radix(x) base b of integers

range(x) decimal exponent range

tiny(x) smallest positive value (real)

Note that all of these functions are generic, and can be used with any supported KIND.

Numeric Manipulation Functions

Using the same representational model as the inquiry functions. Designed to return values related to components of real type.

exponent(x)	value of <i>e</i> in real model
fraction(x)	fractional part in real model
nearest(x,s)	value nearest x in direction of sign of s
rrspacing(x)	reciprocal of relative spacing, $ xb^{-e} b^p$
scale(x,i)	xb ⁱ
set_exponent(x,i)	xb^{i-e}
spacing(x)	b^{e-p} if $x/=0$ and in range, else TINY

IEEE Arithmetic in Fortran

The Fortran 2003 (and later) standard contains facilities for IEEE exception handling. The IEEE¹/ ISO² standard for floating-point arithmetic greatly helped in developing portable numeric code. The goal is to allow for a portable way to test and set the five floating-point exception flags in the IEEE standard.

¹IEEE 754-1985, Standard for floating-point arithmetic

²IEC 559:1989, Binary floating-point arithmetic for microprocessor systems

IEEE Exceptions

IEEE exceptions:

Overflow the result of an operation exceeds the data format Division by Zero finite numerator, zero denominator (result is $\pm \infty$) Invalid operation invalid (e.g. $\infty \times 0$) - result is NaN Underflow result of operation too small for data representation Inexact result of operation can not be represented without rounding

IEEE Intrinsics

Three intrinsic modules are provided:

- IEEE EXCEPTIONS
- IEEE_ARITHMETIC, itself loads IEEE_EXCEPTIONS
- IEEE_FEATURES

Inability to load one of these modules will indicate a non-compliant compiler. For detailed usage, see, for example, Metcalf & Reid. At this point there are relatively few compilers that have explicit support for these modules.

Modules for OO Programming

Note that Fortran modules can be used as objected oriented programming (OOP) constructs:

- Creation of derived types that behave just like intrinsic types
- Intrinic types and operators can be overloaded
- Data can be hidden (encapsulated)
- In such a way one can create semantic extensions

Generic Interfaces

User-supplied functions, like most of the intrinsics, can have **generic** interfaces. For example, consider the intrinsic ABS(x) - behind the scenes, the function actually called depends on the KIND of the argument:

```
CABS for x complex
ABS for x real
IABS for x integer
```

Generic Interface Procedure Example

You can also make use of generic interfaces, of course:

```
MODULE useful 1
     IMPLICIT NONE
     INTERFACE exemplify
      MODULE PROCEDURE exemplify_int    ! Fortran 95 allows separated
      MODULE PROCEDURE exemplify real ! module procedure statements
      MODULE PROCEDURE exemplify complex
     END INTERFACE exemplify ! only Fortran >=95 allows specification at END
    CONTAINS
     SUBROUTINE exemplify int(x)
10
      integer, dimension(:), intent(inout) :: x
11
12
     END SUBROUTINE exemplify real(x)
13
     SUBROUTINE exemplify_real(x)
14
      real, dimension(:), intent(inout) :: x
15
16
     END SUBROUTINE exemplify_complex(x)
17
     SUBROUTINE exemplify complex(x)
18
      complex, dimension(:), intent(inout) :: x
19
20
     END SUBROUTINE exemplify complex(x)
21
    END MODULE useful 1
```

Generic Interface Example (cont'd)

```
1 PROGRAM Main
2 IMPLICIT NONE
3 USE useful_1
4 real :: rx(1000)
5 integer :: ix(1000)
6
7 :
8 CALL exemplify(rx) ! generic call
9 CALL exemplify(ix) ! generic call
10 :
END PROGRAM Main
```

The rule being that the argument list makes the actual choice of routine unambiguous.

Operator Overloading

Operators can be overloaded using the INTERFACE OPERATOR statement:

- Specify the MODULE PROCEDURE/ to deal with the implementation
- Useful (if not required) for derived types
- Operator name/symbox can be any of the intrinsics or any sequence of 31 characters or less in length enclosed in periods (other than .true. and .false)
- Be wary when overloading intrinsic operators binary operators can not be made unary, etc.
- Can not redefine intrinsically defined operations (must remain unambiguous)

Operator Overloading Example

```
MODULE useful 2
     IMPLICIT NONE
     TYPE frac ! type for integer fractions
        integer :: numerator, denominator
     END TYPE frac
6
     INTERFACE OPERATOR (*)
 7
      MODULE PROCEDURE frac int, int frac, frac frac
     END INTERFACE
9
    CONTAINS
10
     FUNCTION frac int(left, right)
11
       type(frac), intent(in) :: left
12
       integer,intent(in) :: right
13
14
     END FUNCTION frac int
15
     FUNCTION int_frac(left, right)
16
       integer,intent(in) :: left
17
      type(frac), intent(in) :: right
18
19
     END FUNCTION int frac
20
     FUNCTION frac frac(left, right)
21
      type(frac), intent(in) :: left
22
      type(frac), intent(in) :: right
23
24
     END FUNCTION frac frac
25
    END MODULE useful 2
```

Note that operator precedence remains unaffected, however.

User Defined Operators

and in a similar way, you can define your own operators:

```
MODULE useful_2
IMPLICIT NONE
INTERPACE OPERATOR (.converged.) ! new op

MODULE PROCEDURE testconv

END INTERFACE

END MODULE PROCEDURE

END WODULE PROCEDURE

END MODULE Useful_2
```

User Defined Assignment

The assignment operator (=) is a little special - overloading it requires an INTERFACE ASSIGNMENT with a SUBROUTINE:

- The first argument has INTENT (OUT) and represents the left-hand side of the assignment
- The second argument has INTENT (IN) and represents the right-hand side
- The subroutine must be "pure," i.e. not alter global data, or produce any output

User Defined Assignment Example

```
MODULE useful 3
     IMPLICIT NONE
     INTERFACE ASSIGNMENT (=)
       MODULE PROCEDURE frac eq
     END INTERFACE
     PRIVATE frac_eq
    CONTAINS
     SUBROUTINE frac_eq(lhs, rhs)
      type (frac), intent (out) :: lhs
10
      type(frac), intent(in) :: rhs
11
       : ! body has to have an assignment to lhs
12
     END SUBROUTINE frac_eq
13
14
    END MODULE useful 3
```

Coarray Fortran Basics

Coarray Fortran (CAF, requires Fortran >=2008) follows the PGAS (partitioned global address space) model. It is composed of simple extensions to "regular" Fortran, similar to UPC (Unified Parallel C) and Titanium (Java).

- Designed to follow SPMD (Single Program Multiple Data)
- Fixed number of processes/threads/images
- Explicit data decomposition and synchronization, data and computing are local
- One-sided communciations through co-dimensions
- Current CAF design is very "minimalist," introduced as few extensions to Fortran as possible

CAF Execution Model

CAF can be used in shared and distributed memory systems (as long as the runtime system supports it),

- image refers to each copy of the program in CAF, with 1 < this image() < num images()
- Co-array syntax uses usual () to refer to local data, [] to refer to remote data
- An image moves remote data to local data through co-array syntax
- Programmer handles synchronization

CAF Coarrays

Data stored in other CAF images are referenced through cosubscripts (enclosed in square brackets), mapped to an image index (one to num_images()).

- Images have their own data, accessed in the standard Fortran way
- Data with codimensions (square brackets) has corank (at decalaration) and cobounds/coextent (at declaration or allocation)
- Coarray data has the same size and shape on each image

sync all

Each CAF image executes on its own without regard to others until image control statements are reached, the simplest of which is sync all, which forces a barrier synchronization.

The above code snippet forms a broadcast, the first sync all ensures that image 1 does not interfere with any other image's use of a, the second that no image uses the old value of a before its update by image one.

More Synchronization

- sync images (image set)
 Synchronizes calling image with all others in the image set sync all is equivalent to sync images (*)
- lock/unlock
 Requires scalar lock variable of type lock_type defined by intrinsic module iso_fortran_env
- critical/end critical restricts execution to one image at a time
- sync memory
 Gives the ability to define boundaries on image between
 segments (statements between two image control statements),
 allows user-defined ordering (very flexible, but also very difficult to
 debug)

All synchronization statements have optional stat= and errmsg= arguments (as with allocate/deallocate).

Coarray Fortran Simple Example

Once again, our favorite "Hello, world" example:

2

4

7

10

11

12

13 14

15

16

Note that Intel's **ifort** has CAF support, and an extra environment variable for controlling the number of images, and it is pretty simple to run in *shared* mode (i.e., on a single node):

```
[rush:~/d fortran/d caf]$ ifort -o hello caf -coarray hello caf2.f90
2
    [rush:~/d fortran/d caf]$ export FOR COARRAY NUM IMAGES=8
    [rush:~/d fortran/d cafl$ ./hello caf2
     Hello, world, I am image 1 of
                                       8 total images on k07n14
     Hello, world, I am image
                               2 of
                                       8 total images on k07n14
     Hello, world, I am image 3 of 8 total images on k07n14
     Hello, world, I am image 4 of 8 total images on k07n14
     Hello, world, I am image 5 of 8 total images on k07n14
     Hello, world, I am image 6 of 8 total images on k07n14
10
     Hello, world, I am image
                             7 of
                                       8 total images on k07n14
11
     Hello, world, I am image
                                8 of
                                       8 total images on k07n14
```

Note that running CAF in shared mode does not require jumping through any extra hoops (not so in the distributed case).

```
#!/bin/bash
     #SBATCH --nodes=2
    #SBATCH --ntasks-per-node=8
     #SBATCH --constraint=CPU-L5520|CPU-L5630
 5
    #SBATCH --partition=debug
 6
     #SBATCH --time=00:10:00
 7
     #SBATCH --mail-type=END
     #SBATCH --mail-user=jonesm@buffalo.edu
9
     #SBATCH --output=slurmO.out
10
     #SBATCH --job-name=caf-test
12
     #export | grep SLURM
13
     module load intel-mpi intel
14
     module list
15
     ifort -o hello caf.dist -coarray=distributed hello caf.f90
16
     EXEC=./hello caf.dist
17
     export MY NODEFILE=tmp.$$
18
     srun -1 hostname -s | sort -n | awk '{print $2}' | uniq > $MY NODEFILE
19
     NNODES='cat $MY NODEFILE | wc -1'
20
     export FOR_COARRAY_CONFIG_FILE=caf.$$
21
     NODES='cat $MY_NODEFILE | awk '{printf "%s ", $1}''
22
     for node in $NODES: do
23
       echo "-n $SLURM NTASKS PER NODE -host $node $EXEC " >> $FOR COARRAY CONFIG FILE
24
     done
25
     #export I MPI DEBUG=4 # enable to see detailed placement info
26
     mpdboot -n $NNODES -f $MY NODEFILE -v
27
     mpdt race
28
     SEXEC
29
       -e $MY_NODEFILE ] && \rm $MY NODEFILE
30
       -e $FOR COARRAY CONFIG FILE | && \rm $FOR COARRAY CONFIG FILE
```

Note the use of mpds and the MPI subsystem for distributing tasks, although the MPI task launcher is not itself being used.

```
2
 3
 4
 5
 6
 7
 8
 9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
```

```
[rush:~/d fortran/d caf]$ cat slurmQ.out
Currently Loaded Modulefiles:
  1) null
                       3) use.own
                                           5) intel-mpi/5.0.1
  2) modules
                      4) intel/15.0
running mpdallexit on d16n32
LAUNCHED mpd on d16n32 via
RUNNING: mpd on d16n32
LAUNCHED mpd on d16n34 via d16n32
RUNNING: mpd on d16n34
d16n32
d16n34
 Hello, world, I am image
                            3 of
                                   16 total images on d16n32
 Hello, world, I am image
                          4 of
                                   16 total images on d16n32
 Hello, world, I am image
                            9 of
                                   16 total images on d16n34
 Hello, world, I am image
                          5 of
                                   16 total images on d16n32
 Hello, world, I am image
                           10 of
                                   16 total images on d16n34
 Hello, world, I am image
                         6 of
                                   16 total images on d16n32
 Hello, world, I am image
                           11 of
                                   16 total images on d16n34
 Hello, world, I am image
                         7 of
                                   16 total images on d16n32
 Hello, world, I am image
                           12 of
                                   16 total images on d16n34
 Hello, world, I am image
                           8 of
                                   16 total images on d16n32
 Hello, world, I am image
                           13 of
                                   16 total images on d16n34
 Hello, world, I am image
                          1 of
                                   16 total images on d16n32
 Hello, world, I am image
                           14 of
                                   16 total images on d16n34
 Hello, world, I am image
                          2 of
                                   16 total images on d16n32
                           15 of
 Hello, world, I am image
                                   16 total images on d16n34
 Hello, world, I am image
                           16 of
                                   16 total images on d16n34
```

Example 1, Build/Use Strings and Internal Files

It is fairly common to want to convert to a string, in C/C++ you can cast or use sprintf, and in Fortran you can use an internal file to write to a string.

In the following example we use an internal file (basically a string) to convert an integer to a string, and then build a file name incorporating that string (suppose, for example, that you wanted to write out a separate file for each rank in an MPI application).

```
1
 2
 3
 5
 6
 7
 8
 9
10
11
12
13
14
15
16
17
18
```

```
program testchar
  implicit none
  ! convert integer to string for use in file name
 character(len=*).parameter :: alphabet1='abcdefghijklm'. &
                              alphabet2='nopgrstuvwxyz'
 character(len=*),parameter :: numerals='0123456789'
 character(len=12) :: cstring
  integer :: testint
 print*, 'First four letters: ',alphabet1(1:4)
 print*, 'alphanumeric: ',alphabet1//alphabet2//numerals
  ! test convert int to string
 testint=123
 write (cstring, '(i12)') testint
 print*, " test string = ",TRIM(ADJUSTL(cstring)),"."
 print*, " filename test = ","outfile "//TRIM(ADJUSTL(cstring))//".dat"
end program testchar
```

```
[rush:~/d_fortran]$ ifort -o testchar testchar.f90
[rush:~/d_fortran]$ ./testchar
First four letters: abcd
alphanumeric: abcdefghijklmnopqrstuvwxyz0123456789
test string = 123.
filename test = outfile 123.dat
```

Example 2, Command Line Arguments

New to Fortran >=2003, command line arguments are now supported. The following example illustrates the use of get_command, command_argument_count, and get_command_argument to process command line arguments to a Fortran program.

```
! cmdline.f90 -- simple command-line argument parsing example
! test fortran2003 support for get_command_argument, get_command,
! and command argument count
program cmdline
  implicit none
  character(len=255) :: cmd
  character(len=*), parameter :: version = '1.0'
  character(len=32) :: arg, date*8, time*10, zone*5
  logical :: do time = .false.
  integer :: i
  call get_command(cmd)
  write(*,*) 'Entire command line:'
  write (*,*) trim(cmd)
  do i = 1, command argument count()
     call get command argument (i. arg)
     select case (arg)
     case ('-v', '--version')
        print '(2a)', 'cmdline version ', version
        stop
     case ('-h', '--help')
        call print_help()
        stop
     case ('-t', '--time')
        do time = .true.
     case default
        print '(a,a,/)', 'Unrecognized command-line option: ', arg
        call print help()
        stop
     end select
  end do
```

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```

```
! Print the date and, optionally, the time
 call date and time(DATE=date, TIME=time, ZONE=zone)
 write (*, '(a,"-",a,"-",a)', advance='no') date(1:4), date(5:6), date(7:8)
 if (do time) then
    write (*, '(x,a,":",a,x,a)') time(1:2), time(3:4), zone
 else
    write (*, '(a)') ''
 end if
contains
 subroutine print_help()
   print '(a)', 'usage: cmdline [OPTIONS]'
   print '(a)', ''
   print '(a)', 'Without further options, cmdline prints the date and exits.'
   print '(a)', ''
   print '(a)', 'cmdline options:'
   print '(a)', ''
   print '(a)', ' -v, --version print version information and exit'
   print '(a)', ' -t, --time
                               print time'
 end subroutine print help
end program cmdline
```

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```

```
[rush:~/d fortran]$ ifort -o cmdline cmdline.f90
 [rush:~/d fortran]$ ./cmdline
 Entire command line:
  /cmdline
2014-09-29
[rush:~/d fortran]$ ./cmdline -v
 Entire command line.
  ./cmdline -v
 cmdline version 1.0
[rush:~/d fortran]$ ./cmdline --help
 Entire command line:
./cmdline --help
usage: cmdline [OPTIONS]
Without further options, cmdline prints the date and exits.
 cmdline options:
  -v, --version print version information and exit
                  print usage information and exit
  -h, --help
  -t, --time
                  print time
[rush:~/d fortran]$ ./cmdline --time
Entire command line:
  ./cmdline --time
2014-09-29 14:13 -0400
```