FEAP - - A Finite Element Analysis Program

Version 8.2 Programmer Manual

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Chapter 1

INTRODUCTION

In this part of the FEAP manual some of the options to extend the capabilities of the program are described. We begin by describing the utilities provided in FEAP for use in data input. Options to add user commands for mesh and command language extensions is then described and finally the method to add an element to the program is described.

1.1 Setting Program Options

The size of problems which may be solved by *FEAP* depends on the amount of memory available in the computer, as well as, solution options used. Memory for the main arrays used to solve problems is dynamically allocated during the solution. Arrays are allocated and deallocated using a system subprogram PALLOC or, for user developed modules using subprogram UALLOC. Further information on use of these routines is given in Section 3.

The IPR parameter in the FEAP82 module controls the specification of the size of REAL variables. For typical UNIX and PC systems all real variables should be double precision and IPR is set to 2. For systems in which REAL*8 variables are *single precision* with the same working length as integer variables the IPR parameter is set to 1. Any error in setting this parameter may lead to incorrect behavior of the program, consequently, do not reset the parameter to single precision unless a careful assessment of compiler behavior for REAL*8 variables has been made.

By placing an alphanumeric version of each manual page in a separate file which has the name of the command and a .t extender (e.g. coor.t for the mesh coordinate input command) it is possible to read each page during execution using the HELP, name command (where name is the command name whose manual page is to be read). For this option to work properly it is necessary to define the path name to each manual page in the FEAP82 module. For example:

file(1) = 'c:\Feap\Manual\Mesh\'
file(2) = 'c:\Feap\Manual\Macr\'
file(3) = 'c:\Feap\Manual\Plot\'

defines a typical path for a PC system. Each system requires a proper path definition. FEAP will add the requested command name to each of the above paths to find mesh, solution, or plot commands.

Normally *FEAP* reads each input data line as text data and checks each character for the presence of parameters, expressions, and constants. For very large data sets this parsing of each instruction can consume several seconds of compute time. If all data is normally provided as numerical data, without use of any parameters or expressions, the input time may be reduced by setting the value of the logical variable COFLG in FEAP82 to *false*. *FEAP* will automatically switch to parsing mode if any record contains non-numerical data item. It is also possible to use the PARSe and NOPArse commands to set the appropriate mode of data input.

During the input of plot commands *FEAP* has the option to either set input options automatically (DEFAult mode) or to read the values or range of contours to plot. The default mode of operation may be assigned in the FEAP82 module by setting the variables DEFALT and PROMPT. Setting DEFALT to *true* indicates that all default options are to be set automatically. If DEFALT is set *false*, a prompt for contour intervals may be requested by setting PROMPT to *true*.

FEAP has options to produce encapsulated PostScript output files in either gray scale of in color. The default mode may be established by setting the variable PSCOLR and PSREVS. Setting PSCOLR true indicates the PostScript files will be in color (unless set otherwise by the PLOT COLOr data command. The PSREVS variable reverses the color sequence.

Arrays in *FEAP* are dynamically allocated during execution. Thus, it is possible to define and destroy arrays as well as to increase or decrease the size of an array. A parameter is provided to control when an array is to be decreased in size. The parameter is **INCRED** and an array is decreased in size only when the new size is less than the old size by the assigned value.

The last parameter which may be set in the FEAP82 module is the level for displaying available commands when the HELP command is used while in mesh, solution, or plot

mode. *FEAP* contains a large number of commands which are not commonly used by many users. To control the default number of commands displayed to users the commands have been separated into four levels: (0) Basic; (1) Intermediate; (2) Advanced; and (3) Expert. The level to be displayed when using the HELP command is given may be set in the integer variable HLPLEV. That is, setting:

hlplev = 1 ! Intermediate

results in commands up to the *intermediate* level being displayed. It is possible to raise or lower the level during execution using the command MANUal,,level where level is the numerical value desired.

When developing program modules it is often desirable to have output of specific quantities available (e.g. tracking the change in some parameters during successive iterations. *FEAP* provides for a switch to make the outputs active or inactive during an execution. The switch is named **debug** and placed in

integen		ndebug	
logical	L		debug
common	/debugs/	ndebug	debug

The value of the debug is set true by the solution command DEBUg and false by the command DEBUg, OFF. Thus, placing code fragments into modules as

```
if(debug) then
    write(iow,*) 'LABEL',list ... ! writes to output file
c and/or
    write( *,*) 'LABEL',list ... ! writes to screen
    endif ! debug
```

This device supplements use of available debuggers on the computer.

1.2 Uses of Common and Include Statements

FEAP contains many COMMON statements that are used to pass parameters and small array values between subprograms. For example, access to the debugging parameter **debug** is facilitated through **common /debugs/**. Users may either place the common statement (as well as data typing statements) directly in the routine or may use an include statement. For debugging the statement would be

include 'debugs.h'

which during compilation would direct the precompiler to load the current common statement from this file. In *FEAP* all include files have the same name as the common with an added extender .h. The only exception is a dummy blank common which uses the file name comblk.h which is defined as

real*8 hr integer mr common hr(1),mr(1)

The dummy arrays hr(1) and mr(1) serve to pass all dynamically allocated arrays between subprograms using a pointer array contained in the common array named np(*) [or for user defined arrays in up(*)] located in the include file pointer.h. See Section 3 for more details on use of pointers. All include files are located in the directories include and include/integer4 or include/integer8. The subdirectory integer4 defines 32-bit integer parameters and integer8 defines 64-bit integer parameters.

It is highly recommended that users use include files rather than giving equivalent common statements directly. If later releases of the *FEAP* program revise contents in a common block, it will only be necessary to recompile the user routine rather than change all the common statement definitions. Also, by defining the correct path in the **makefile.in** or your compiler it is not necessary to modify any routine when switch from 32-bit machines to 64-bit machines.

Chapter 2

DATA INPUT AND OUTPUT

FEAP includes utilities to perform input and to output small arrays of data. Users are strongly encouraged to use the input utilities but often may wish to use their own utilities to output data.

2.1 Parameters and Expressions

The subroutines PINPUT and TINPUT are input subprograms used by FEAP to input each data record. They permit the data to be in a free form format with up to 255 characters on each record, as well as to employ expressions, parameters, and numerical representations for each data item. These routines also should be used to input data in any new program module developed. The PINPUT routine returns data to the calling subprogram in a double precision array. The following statements may be included as part of the routine performing the input.

```
subroutine xxx(....)
logical errck, pinput
integer ior,iow,ilg
common /iofile /ior,iow,ilg
real*8 td(5)
1 if(ior.lt.0) write(*,3000)
errck = pinput(td, 5)
if(errck) go to 1
```

The parameters defined in the common block are:

ior	- input file unit number (if negative, input	j
	from keyboard)	
iow	- output file unit number	
ilg	- solution log file unit number	

If an error occurs during input from the keyboard *FEAP* returns a value of true for the function and a user may reinput the record if the implied loop shown above is used. For inputs from a file, the program will stop and an error message indicating the type of error occurring and the location in an input file is written to the output file.

The input routines return data in a real*8 array td(*). If any td(i) is to be used as an integer or real*4 quantity, it must be cast to the correct type. That is, the following operations should be used to properly cast the variable type:

```
real*4 t
real*8 td(5)
integer j
logical errck, pinput
errck = pinput (td, 5)
j = nint( td(1)) ! Integer assignment
t = float(td(2)) ! Real*4 assignment
```

PINPUT may be used to input up to 16 individual expressions on one input record (each input record is, however, limited to 255 characters).

The routine TINPUT differs from PINPUT by permitting text data to also be input. It is useful for writing user commands or to input data described by character arrays. The routine is used as

```
logical errck, tinput
integer nt, nn
character text(16)*16
real*8 td(16)
errck = tinput(text,nt,td,nn)
```

The parameter **nt** specifies the number of *text* values to input and the **nn** specifies the number of *real data* values to input. The value for parameter **nt** or **nn** may be zero. Thus the use of

errck = tinput(text,0,td,nn)

is equivalent to

```
errck = pinput(td,nn)
```

Text variables may be converted to numerical (REAL*8) form using the subroutine call

call setval(text,nc,td)

where text is a string with nc characters and td a REAL*8 variable. The text string can contain any parameters, expressions or numerical constants which evaluate to a *single* value.

2.2 Array Outputs

Two subprograms exist to output arrays of integer and real (double precision) data. The routine MPRINT is used to output real data and is accessed by the statement:

call mprint(array, nrow, ncol, ndim, label)

where **array** is the name of the array to print, **nrow** and ncol are the number of rows and columns to output, **ndim** is the first dimension on the array, and **label** is a character label which is added to the output. For example the statements:

```
real*8 aa(8,6)
. . .
    call mprint( aa(2,4), 2, 3, 8, 'AA')
```

outputs a 2×3 submatrix from the array **aa** starting with the entry **aa(2,4)**. The output entries will be ordered as the terms:

aa(2,4) aa(2,5) aa(2,6) aa(3,4) aa(3,5) aa(3,6)

The MPRINT routine adds row and column labels as well as the character label.

The routine **IPRINT** is used to output integer data and is accessed by the statement:

call iprint(array, nrow, ncol, ndim, label)

where all parameters are identical to those for MPRINT except the array must be of type integer.

Chapter 3

ALLOCATING ARRAYS

Dynamic data allocation is accomplished in *FEAP* by defining addresses in pointers contained in the common block defined in pointer.h. This common block contains pointers np for standard program arrays and up for user defined arrays and has the form

integer num_nps , num_ups
parameter (num_nps = 400 , num_ups = 200)
integer np , up
common /pointer/ np(num_nps) , up(num_ups)

Each pointer is an offset relative to the address of a REAL*8 array hr(1) or an INTEGER array mr(1) defined in a blank common

real*8 hr integer mr common hr(1),mr(1)

which is placed in the file comblk.h in the include/integer4 or include/integer8 directory. The pointers in the integer4 subdirectory have 32-bit lengths and should be used for cases where addressing is within 4 GByte range. The pointers in the integer8 subdirectory have 64-bit lengths and should be used for additional index ranges. The arrays 'hr' and 'mr' are used to establish addresses only and not to physically store data. This mechanism permits references to elements in arrays which have positions relative to hr or mr that may be after or before 1. Thus, FEAP must be compiled without strict array bound checking. Size of problems is limited only by the available memory in the computer used.

When using 64-bit pointers users must be careful to always define the address of an array in a calling statement to also be 64-bits in length. For example use of

```
integer ioff
...
ioff = np(111) + numnp
call submat( hr(ioff), ...)
```

would cause an error since the pointer ioff is only 32 bits in length. To avoid this problem it is necessary to either declare ioff to be 64-bits long as

```
integer*8 ioff
```

or use one of the *FEAP* include files p_int.h (defining the integer type array fp(10)) or p_point.h (defining the integer type scalar point). A type of correct length is controlled at compile time by which subdirectory is used (i.e., integer4 or integer8).

Using this scheme permits direct reference to either real*8 or integer arrays in program modules without need to pass arrays through arguments of subprograms. A subprogram PALLOC controls the allocation of all standard arrays in *FEAP* defined by the np pointers and a subprogram UALLOC permits users to add allocation for their own arrays defined by the pointers up. The basic use of the routines is provided by an instruction

```
setvar = palloc(number,'NAME',length,precision)
```

or

```
setvar = ualloc(number,'NAME',length,precision)
```

where setvar, palloc and ualloc are logical types, number is an integer number of the array, NAME is a 5 character name of the array, length is the number of words of storage needed for the array, and precision is the type of array to allocate (1 for integer and 2 for real*8 types). Upon initial assignment of any array its values are set to zero. Thus, if the array is to be used only once it need not be set to zero before accumulating additional values. If the array is to be reused or resized (see below) it must be reinitialized prior to accumulating any additional values. Use of these subprograms controls the assignment of memory space for all arrays such that no conflicts occur between hr and mr referenced arrays. Each routine which makes direct reference to an allocated array using a pointer (e.g., hr(np(43)) or mr(up(1))) must contain include files as

include 'pointer.h'
include 'comblk.h'

As an example for the use of the above allocation scheme consider a case where it is desired to allocate a real (double precision array) with length NUMNP (number of nodes in mesh) and an integer array with length NUMEL (number of elements in mesh). The parameters NUMNP and NUMEL are contained in COMMON /CDATA/ and available using the include file cdata.h. The new arrays aredefined using the temporary names TEMP1 and TEMP2 which have numerical locations '111' and '112', respectively.¹ The two arrays are allocated using the statements

setvar = palloc(111, 'TEMP1', numnp, 2)
setvar = palloc(112, 'TEMP2', numel, 1)

where the last entry indicates whether the array is REAL*8 (2) or INTEGER (1). These arrays are now available in any subprogram by specifying the pointer.h and comblk.h include files and referencing the arrays using their pointers, e.g., in a subroutine call as:

```
include 'pointer.h'
include 'comblk.h'
...
call subname ( hr(np(111)) , mr(np(112)) .... )
```

Note the use of hr(*) and mr(*) for the double precision and integer references, respectively. Also, the use of the pointers avoids a need to include the array reference until it is needed in a computation.

A short list of the mesh arrays available in *FEAP* is given in Table 3.1, for solution arrays in Table 3.2, and for element arrays in Table 3.3. The names of all active arrays in any analysis may be obtained using the SHOW, DICTionary solution command. Tables 3.4 and 3.5 describe the use of individual entries in the arrays IX and IE, respectively.

¹See the subprogram palloc.f in the program directory for the names and numbers of existing arrays.

NAME	Num.	dim 1	dim 2	dim 3	Description
ANG	45	numnp	-	-	Angle
D	25	ndd	nummat	-	Material parameters
F	27	ndf	numnp	2	Force and Displacement
ID	31	ndf	numnp	2	Equation nos.
IE	32	nie	nummat	-	Element control, dofs, etc.
IX	33	nen1	numel	-	Element connections
Т	38	numnp	-	-	Temperature
U	40	ndf	numnp	3	Solution array
VEL	42	ndf	numnp	nt	Solution rate array
Х	43	ndm	numnp	-	Coordinates

Table 3.1: Mesh Array Names, Numbers and Sizes

NAME	Num.	dim 1	dim 2	dim 3	Description
CMASn	n+8	compro	-	-	Consistent Mass
DAMPn	n+16	compro	-	-	Damping
JPn	n+20	neq	-	-	Profile pointer
LMASn	n+12	neq	-	-	Lump Mass
TANGn	n	maxpro	-	-	Symmetric tangent
UTANn	n+4	maxpro	-	-	Unsymmetric tangent

Table 3.2: Solution Array Names, Numbers and Sized

NAME	Num.	dim 1	dim 2	dim 3	Description
ANGL	46	nen	-	-	Angle
LD	35	nst	-	-	Assembly nos.
Р	35	nst	-	-	Element vector
S	36	nst	nst	-	Element matrix
TL	39	nen	-	-	Temperature
UL	41	ndf	nen	6	Solution array
XL	44	ndm	nen	-	Coordinates

Table 3.3: Element Array Names, Numbers and Sizes

NAME	Description
IX(1 ,e)	Global node 1
	to
IX(nen ,e)	Global node nen
IX(nen+1,e)	H1 history data pointer
IX(nen+2,e)	H2 history data pointer
IX(nen+3,e)	H3 history data pointer
IX(nen1 ,e)	Element material type number
IX(nen1-1,e)	Element region number (default $= 0$)
IX(nen1-2,e)	Active/deactive indicator
IX(nen1-3,e)	Active/deactive start

Table 3.4: Element connection array IX use for element e

NAME	Description
IE(1,ma)	Global DOF-1
	to
IE(ndf,ma)	Global DOF-ndf
IE(nie ,ma)	Number history variables/element (NH1 and NH2)
IE(nie-1,ma)	Element material type number (ELMT01 = 1, etc.)
IE(nie-2,ma)	Element material type identifier (default $=$ ma)
IE(nie-3,ma)	Offset to $NH1/2$ history variables (default = 0)
IE(nie-4,ma)	Offset to NH3 history variables (default $= 0$)
IE(nie-5,ma)	Number history variables/element (NH3)
IE(nie-6,ma)	Finite rotation update number (for UROTxx)
IE(nie-7,ma)	Get tangent from element if 0; if > 0 numerically differ-
	entiate residual to obtain tangent.
IE(nie-8,ma)	Equation number for element Lagrange multiplier

Table 3.5: Element control array IE use for material number ma

The subprograms PALLOC and UALLOC may also be used to destroy a previously defined array. This is achieved when the length of the array is specified as zero (0). For example, to destroy the arrays defined as TEMP1 and TEMP2 the statements

```
setvar = palloc( 111, 'TEMP1', 0, 2 )
setvar = palloc( 112, 'TEMP2', 0, 1 )
```

are given. Use of these statements results in the pointers np(111) and np(112) being set to zero and the space used by the arrays being released for use by other allocations at a later point in the program.

A call to PALLOC or UALLOC for any previously defined array but with a different nonzero length causes the size of the array to be either increased or decreased.

For user defined arrays specified in UALLOC care should be exercised in selecting the alphanumeric NAME parameter, which is limited to 5 characters, so that conflicts are not created with existing names (use of the SHOW, DICT command is one way to investigate names of arrays used in an analysis) or check the names already contained in the subprogram PALLOC.

The subroutine PGETD also may be used to retrieve internal data arrays by NAME for use in user developed modules. For example, if a development requires the nodal coordinate data the call

```
integer xpoint, xlen, xpre
logical flag
....
call pgetd ('X ',xpoint,xlen,xpre,flag)
```

will return the first word address in memory for the coordinates as xpoint, the length of the array as xlen, and the precision of the array as xpre. If the retrieval is successful flag is returned as true, whereas if the array is not found it is false. The precision will be either one (1) or two (2) for INTEGER or double precision (REAL*8) quantities, respectively. Thus, the above coordinate call will return xpre as 2 and xlen will be the product of the space dimension of the mesh and the total number of nodes in the mesh. The first coordinate, x_1 , may be given as

x1 = hr(xpoint)

any other coordinates at nodes may also be recovered by a correct positioning in later words of hr. For example y_1 is located at hr(xpoint+1). The use of pgetd can lead

to errors for situations in which the length of arrays changes during execution, since in these cases the value of the pointer xpoint can change. For such cases a call to pgetd must be made prior to each reference involving xpoint. On the other hand, reference using the pointers defined in arrays NP or UP are adjusted each time an array changes size. However, users must ensure that a calling sequence is not sensitive to a change in pointer. One way pointer changes can still lead to errors is through a program

```
call subname ( hr(np(111)), mr(np(112)), ....)
```

and then change the length of the array number '111' or '112' in the subroutine.

Chapter 4

USER FUNCTIONS

Users may add their own procedures to facilitate additional mesh input features, to perform transformations or manipulations on mesh data, to add new solution commands, or to add new plot capabilities.

4.1 Mesh Input Functions - UMESHn.

To add a mesh input command a subprogram with the name UMESHn, where n has a value between 0 and 9 must be written, compiled, and linked with the program. The basic structure of the routine UMESH1 is:

```
subroutine umesh1(uprt)
С
      User defined routine to input mesh data to FEAP
      implicit
               none
                          ! Contains UCT variable
      include
               'umac1.h'
      logical
                uprt
с
      Set name 'mes1' to user defined
      if(pcomp(uct,'mes1',4)) then
                       ! Set user defined command name
        uct
            = 'XXXX'
      else
        User execution function statements follow
С
      end if
```

end

The parameter UPRT is a logical parameter which is set to false when the NOPRint mesh command is given and to true when the PRINt command is used (default is true). The common block UMAC1 transfers the character variable UCT for the name of the command. The default names are MESn where n is the same as the routine name number. Assignment of a unique character name (which must not conflict with names already assigned for *mesh input commands*) should be used to replace the xxxx shown.

When *FEAP* begins execution it scans all of the UMESHn routines and replaces the command names mes1, etc., by the user furnished names. Thus, when the command HELP is issued while in interactive MESH mode, the user name will appear in the list instead of the default name (note, *FEAP* does not always display all available commands. To see all commands issue the command MANUal, 3 and then the HELP command).

The ability to get array names as shown in Chapter 3 can be used to develop user routines for input of coordinates, element connections, etc. With this facility it is possible to develop an ability to directly input data prepared by other programs which may be in a format which is not compatible with the requirements of standard FEAP mesh commands.

4.1.1 Command line data

It is possible to include extra data on the command line for user functions. If the data is not passed as an argument (e.g., as in the UMESHn functions) the data is recovered from a character string yyy*255 contained in the labeled common statement chdata. The string is formatted into words with a field widths of 15 characters. Text words are left justified and numerical values are right justified. The data must be recovered in the function before any additional input statements are processed. For example if a user input function has the command line:

GETData VALUes 35

is developed in the user function UMESH1 the first argument GETData is assigned to the variable uct as the name of the function (see above), however, by default the other two words are ignored. To recover their values the statement

include 'chdata.h'

```
character lct*15
real*8 ctl
integer itl
```

is added to the user function and the items recovered in the **else** option of the function using the statements:

lct = yyy(16:30)
call setval(yyy(31:45),15, ctl)

If *FEAP* parameters or expressions are used to define a numerical entry the subprogram **setval** will make the appropriate evaluation. If necessary, the real value for **ctl** can be cast into an integer using

itl = nint(ctl)

Note that the total number of added words must be 15 items or less (this is imposed by the total of 16 items on any FEAP input record).

4.2 User Material Models

Users may add material models to elements by appending subprograms UMATIn and UMATLn (where n have values from 0 to 9) to the *FEAP* system. The subprogram UMATIn defines parameters used by the model and the subprogram UMATLn is called by the element for *each* computation point (i.e., the quadrature point), receives the value of a deformation measure as input and must return the value of stress and tangent moduli as output.

To activate a user material model the input data for the mesh MATErial command must include a statement with UCON as the first field. For example in a solid element the command sequence can be

```
MATErial ma
SOLId
UCONstitutive xxxx v1 v2 ...
```

The role of the xxxx and vi data will be described in Section 4.2.1.

It is possible to use standard input parameters defined in Tables 5.5 to 5.8, as well as by preceding the UCON command with a normal input sequence. For example, if isotropic elastic properties are needed they may be included in the input sequence as

MATErial ma SOLId ELAStic ISOTropic e nu UCONstitutive xxxx v1 v2 ...

No standard commands should follow the UCON command.

Alternatively, users may input elastic properties as part of their UMATIn module. If the user routine does input additional data records (after the UCON record) and these are terminated by a blank record, a second blank record will be needed to discontinue material data input for this set. In all cases at least one blank record is always needed to terminate the input of standard options for the material set.

4.2.1 The UMATIn Module

A sample module for a user constitutive model is shown in Fig. 4.1. As shown in this figure, the UMATIn module has 5 arguments. The name of the constitutive equation to be described is passed in the first parameter type. The second parameter passes an array (vv(*)) which may be used to define up to 5 parameters for the material model. The example shown above for the UCON includes the type data as xxxx and the array vv(*) values as v1 v2 Users may also provide additional input within the UMATIn module using the routines PINPUT or TINPUT described in Sect. 2.1. The values of user parameters must be saved in the array ud(*) (the third argument of UMATIn). In the current version there are 50 words of double precision values available by default. Additional values may be allocated by assigning a larger value on the control record (first record after the FEAP title record). Each material model is assigned a user material number to the return parameter umat. This number must be a positive integer. Finally, the number of history parameters to be assigned to each computation (quadrature) point must be returned in the parameter n1. Currently, the parameter n3 may be set but is not available to the user material model. Thus, all history variables must be retained in the n1 list. Use of history variables is described later as part of the UMODEL module.

```
subroutine umati1(type,vv, ud,n1,n3)
c-----[--.--+----.--+-----]
      Purpose: User material model interface
С
с
      Inputs:
                - Name of constitutive model (character variable)
         type
С
с
         vv(*)
                - Parameters: user parameters from command line
С
      Outputs:
                - Number history terms: nh1,nh2
        n1
С
        n3
               - Number history terms: nh3
С
        ud(*) - User material parameters
С
c-----[--.--+----.---+-----]
     implicit none
     include 'iofile.h'
     logical pcomp
     character type*15
              n1,n3
     integer
     real*8
              vv(5), ud(*)
     Specify type of user model
С
     if(pcomp(type,'mat1',4)) then
       type = 'E-1d'
                                  ! Specify new name for model
     Input/output user data and save in ud(*) array
С
       Set values of 'n1' if required
С
       n1 = ...
       write(iow,*) ' User Constitutive Inputs: E = ',vv(1)
       ud(1) = vv(1)
     endif
     end
```

Figure 4.1: Sample UMATI1 module

4.2.2 The UMATLn Module

A sample for the UMATL1 module is shown in Fig. 4.2. This subprogram will be called by many of the elements included within *FEAP* if a user model has been specified as part of the MATE mesh data (see previous subsection). The user model will not be called for truss, frame, plate, and shell elements which use resultant models to describe behavior. Also, any form which requires a one-dimensional model will not use a UMATLn module. The module is designed to compute three-dimensional constitutive models in which the stress and strain are stored as 6-component vectors and the tangent moduli as a 6×6 matrix. Strains are passed to UMATLn in the argument array eps(6) and stored in the order

$$\boldsymbol{\epsilon} = \begin{bmatrix} \epsilon_{11} & \epsilon_{22} & \epsilon_{33} & \gamma_{12} & \gamma_{23} & \gamma_{31} \end{bmatrix}^T$$

where $\gamma_{ij} = 2 \epsilon_{ij}$ is the engineering shearing strain. Stress and moduli are to be associated with the same ordering and returned in the argument arrays dimensioned as sig(6) and dd(6,6), respectively. All values are to double precision (i.e., REAL*8).

When UMATLn is called the model n will be that which is defined in the module UMATIn. Current values of the strains are, as mentioned above, passed in the array eps(6) and the trace of the strain in the parameter theta. Thus,

$$\theta = \epsilon_{ii} = \epsilon_{11} + \epsilon_{22} + \epsilon_{33} \; .$$

In addition, if thermal problems are being solved the current value for the temperature is passed as td. All material parameters for the current model are passed in the arrays d(*) and ud(*). The array d(*) contains parameters assigned by standard *FEAP* commands as described in Tables 5.5 to 5.8 and the array ud(*) contains values as assigned in the user module UMATIn.

For constitutive equations with additional (internal) variables that evolve in *time*, users must define entries for the h1(*) array. The number of entries available in the array for *each evaluation* (i.e., each quadrature point) is nh. The value for nh is defined by the parameter n1 in module UMATIn (see Fig. 4.1). Values from the previous time step are passed back to the module in the array hn(*) (which also contains nh entries). Users should never modify entries in the hn(*) array. Finally, the values of the element operation switch is passed as the parameter isw (See Chapter 5 for operations performed during different values of isw).

Using the above information users *must* compute values for the stress and the associated tangent matrix. These are returned to the element in the arrays sig(6) and dd(6,6). In addition, updates for any of the history parameters must be assigned in the array

h1(*) and returned to the element. Values of history variables returned are not used for all values of isw (e.g., when reporting or projecting stresses under isw = 4 and isw = 8 they are not saved). Values retained in the h1(*) array are copied to the hn(*) array each time the command statement TIME is issued in a solution.

4.2.3 Auto time step control

The solution command:

initiates an attempt to control the solution process by a variable time stepping algorithm based on a user set value in the material constitution. The value to be set is named **rmeas** which is passed between constitution and solution modules in the labeled common

> real*8 rmeas,rvalu logical aratfl common /elauto/ rmeas,rvalu(3),aratfl

The three parameters may be used in defining an acceptable value for **rmeas**. The algorithm coded monitors the solution during a standard iteration process set by, for example:

If during any iteration up to n the value of rmeas exceeds a value of 2 (rmeas = 0 at the start of the loop) a new value of Δt is immediately set to

$$\Delta t_{new} = 0.85 \, \Delta t / rmeas$$

and the iteration process is started over. On the other hand if convergence occurs during the time step and the value of **rmeas** is smaller than 1.25, the time step is adjusted according to

$$\begin{array}{ll} \Delta t_{new} = 1.50 \, \Delta t & ; \ rmeas \leq 0.5 \\ \Delta t_{new} = 1.25 \, \Delta t & ; \ 0.5 < rmeas \leq 0.8 \\ \Delta t_{new} = \Delta t / rmeas & ; \ 0.8 < rmeas \end{array}$$

```
subroutine umatl1(eps,theta,td,d,ud,hn,h1,nh,istrt,sig,dd,isw)
Purpose: User Constitutive Model
С
С
     Input:
         eps(*) - Current strains at point (small deformation)
С
С
                - Deformation gradient at point(finite deformation)
         theta - Trace of strain at point
С
                - Determinant of deformation gradient
С
        td - Temperature change
d(*) - Program material parameters
С
С
         ud(*) - User material parameters
hn(nh) - History terms at point: t_n
С
С
         h1(nh) - History terms at point: t_n+1
С
         nh - Number of history terms
С
         isw - Solution option from element
С
     Output:
С
         sig(6) - Stresses at point.
С
         dd(6,6) - Current material tangent moduli
С
implicit none
     integer umat, nh, isw, i
     real*8
            td
             eps(*),theta(*),d(*),ud(*),hn(nh),h1(nh), sig(6),dd(6,6)
     real*8
     Dummy model: sig = ud(1)*eps
С
       do i = 1, 6
        dd(i,i) = ud(1)
        sig(i) = ud(1) * eps(i)
       end do
     endif
     end
```

Figure 4.2: Sample UMATLn module

Finally, if convergence does not occur with in the n steps, then the time step is reset according to

$$\Delta t_{new} = 0.85 \,\Delta t/rmeas \quad ; \quad 1.25 < rmeas$$
$$\Delta t_{new} = \Delta t/3 \qquad ; \quad \text{otherwise.}$$

After any of the above adjustments the value of rmeas is reset to zero (0).

An optimal value of **rmeas** is 1.25 – which leaves the step unchanged. The above algorithm was proposed by Weber *et al.* [1].

4.3 Mesh Manipulation Functions - UMANIn.

The UMANIn modules, where n ranges from 0 to 9, may be used to perform transformations or manipulations on previously prescribed data. These commands appear between the mesh input END command and the first INTEractive or BATCh solution command. To add a mesh manipulation command a subprogram with the name UMANIn, where n has a value between 0 and 9 must be written, compiled, and linked with the program. The basic structure of the routine UMANI1 is:

```
subroutine umani1(prtu)
      User defined routine to manipulate mesh data for FEAP
С
      implicit none
      include
                'umac1.h'
                           ! Contains UCT variable
      logical
                prtu
      Set name 'man1' to user defined
С
      if(pcomp(uct, 'man1',4)) then
            = 'XXXX'
                          ! Set user defined command name
        uct
      else
        User execution function statements follow
С
      end if
      end
```

The parameter PRTU is a logical parameter which is set to false when the NOPRint mesh command is given and to true when the PRINt command is used (default is true). The common block UMAC1 transfers the character variable UCT for the name of

the command. The default names are MANn where n is the same as the routine name number. Assignment of a unique character name (which must not conflict with names already assigned for *mesh input commands*) should be used to replace the xxxx shown.

After *FEAP* completes the input of mesh data it scans all of the UMANIn routines and replaces the command names man1, etc., by the user furnished names.

The ability to get array names as shown in Chapter 3 can be used to develop user routines for manipulation of the mesh data. For example, if a user has added the specification of information by coordinates it may later be necessary to associate the data with specific node numbers. This can be accomplished using a manipulation command which searches for the node number whose coordinates are closest to the specified location.

4.4 Solution Command Functions - UMACRn.

In a similar manner, users may add solution commands to the program by adding a routine with the name UMACRn where n ranges from 0 to 9.

```
subroutine umacr0(lct,ctl,uprt)
      User solution command function
С
      implicit none
      include
               'umac1.h'
                              ! Contains the variable UCT
      logical
                uprt
      character lct*15
      real*8
                ct1(3)
      Set command word
С
      if(pcomp(uct, 'mac0', 4)) then
        uct = 'xxxx'
      else
        User command statements are placed here
С
      endif
      end
```

The parameters LCT and CTL are used to pass the second word of a solution command and the three parameter values read, respectively. Again the name xxxx should be selected to not conflict with existing solution command names and will appear whenever HELP is issued.

4.5 Plot Command Functions - UPLOTn.

In a similar manner, users may add new plot commands to the program by adding a routine with the name UPLOTn where n ranges from 0 to 9.

```
subroutine uplot0(ctl,uprt)
С
      User plot command function
      implicit none
                              ! Contains the variable UCT
      include
                'umac1.h'
      logical
                uprt
      real*8
                ct1(3)
      Set command word
С
      if(pcomp(uct, 'plt0',4)) then
        uct = 'xxxx'
      else
С
        User plot command statements are placed here
      endif
      end
```

The parameters CTL(3) are used to pass the three parameter values read, respectively. Again the name xxxx should be selected to not conflict with existing plot command names and will appear whenever HELP is issued.

Two plot utilities are available for placing lines on the screen. These are named DPLOT and PLOTL. The calling form for DPLOT is given as

call dplot(s1,s2,ipen)

where s1, s2 are screen coordinates ranging from 0 to 1. Similarly, the calling sequence for PLOTL is

Number	Color
0	Black
1	White
2	Red
3	Green
4	Blue
5	Yellow
6	Cyan
7	Magenta
8	Orange
9	Coral
10	Green-Yellow
11	Wheat
12	Royal Blue
13	Purple
14	Aquamarine
15	Violet-Red
16	Dark Slate Blue
17	Gray
18	Light Gray

 Table 4.1: Color Table for Plots

call plotl(x1,x2,x3,ipen}

where x1, x2, x3 are coordinates values of the mesh. The value of ipen ranges from 1 to 3: 1 starts a filled panel; 2 draws a line from the current previous point to the new point; 3 moves to the new point without drawing a line. If a filled panel is started it must be closed by inserting the statement

call clpan()

Lines are drawn or panels filled in the current color. A color is set using the statement

```
call pppcol(color, switch)
```

where color is an integer defining the color number and switch should be zero. The color values are given in Table 4.1.

Chapter 5

ADDING ELEMENTS

FEAP permits users to add their own element modules to the program by writing a single subprogram called

subroutine elmtnn(d,ul,xl,ix,tl,s,r,ndf,ndm,nst,isw)

where nn may have values between 01 and 50. Each element subprogram must be added before loading the *FEAP* library since dummy subprograms are included in the library to avoid unsatisfied externals. The basic structure for an element routine is shown in Figures 5.1 and 5.1.

Information is provided to the element subprogram through data passed as arguments and data passed in common blocks. The data passed as arguments consists of eleven

subroutine elmtnn(d,ul,xl,ix,tl,s,r,ndf,ndm,nst,isw)
c Prototype FEAP Element Routine: nn = 01 to 50
implicit none
c Common blocks: See Figure 5.2.
integer ndf,ndm,nst,isw
integer ix(nen1,1)
real*8 d(*),ul(ndf,*),xl(ndm,*),tl(*),s(nst,nst),r(nst)
if(isw.eq.0 .and. ior.lt.0) then
c Output element description
write(*,*) ' Elmt 1: Element description'

Figure 5.1: FEAP Element Subprogram. Part 1.

с с с	<pre>elseif(isw.eq.1) then Input/output of property data after command: 'mate' d(*) stores information for each material set Return: nh1 = number of nh1/nh2 words/element Return: nh3 = number of nh3 words/element</pre>
с	elseif(isw.eq.2) then Check element for errors. Negative jacobian, etc.
с с с с с с с с	<pre>elseif(isw.eq.3) then Return: Element coefficient matrix and residual s(nst,nst) element coefficient matrix r(ndf,nen) element residual hr(nh1) history data base: previous time step hr(nh2) history data base: current time step hr(nh3) history data base: time independent</pre>
с с с	<pre>elseif(isw.eq.4) then Output element quantities (e.g., stresses) elseif(isw.eq.5) then Return: Element mass matrix s(nst,nst) consistent matrix r(ndf,nen) diagonal matrix</pre>
c c	elseif(isw.eq.6) then Compute residual only r(ndf,nen) element residual
C C C	<pre>elseif(isw.eq.7) then Return: Surface loading for element s(nst,nst) coefficient matrix r(ndf,nst) nodal forces</pre>
с с с	<pre>elseif(isw.eq.8) then Compute stress projections to nodes (diagonal) p(nen) projection weight: wt(nen) s(nen,*) projection values: st(nen,*) (default: project 8 quantities) endif end</pre>

Figure 5.1: FEAP Element Subprogram. Part 2.

(11) items which are briefly described in Table 5.1^1 .

¹Note in Table 5.1 that *FEAP* transfers the values for most of the solution parameters in array UL(NDF, NEN, *) at time t_{n+a} , where *a* denotes a value between 0 and 1. The value of *a* is 1 (i.e., values are reported for time t_{n+1}) unless generalized midpoint integration methods are used. For the present we will assume *a* is 1.

FEAP carries out tasks according to the value of the parameter ISW passed as the eleventh parameter of the ELMTnn subprogram. A short description of the task carried out by each value, as currently implemented, is shown in Table 5.2.

To use the basic features available in *FEAP* it is necessary to program tasks labeled as R shown above. If elements have local variables that need to be retained between subsequent time steps *history variables* may be defined as described in Section 5.6. In this case it is necessary to code task 12 if special transformations of the variables are required (otherwise merely return with no changes) and if any of the parameters have non-zero initial values task 14 is used to set these values (zero values are set by default). Finally, if special plotting options are desired it may be necessary to program task 20 (note that contours for element variables such as stress, strain, etc. are developed from task 8).

Parameter	Description		
d(*)	Element data parameters		
	(Moduli, body loads, etc.)		
ul(ndf,nen,j)	Element nodal solution parameters		
	nen is number of nodes on an element (max)		
	$j = 1$: Displacement $u_{n+a}^{(k)}$		
	$j = 2$: Increment $u_{n+a}^{(k)} - u_n$		
	j = 2: Increment $u_{n+a}^{(k)} - u_n$ j = 3: Increment $u_{n+1}^{(k)} - u_{n+1}^{(k-1)}$		
	j = 4: Rate $v_{n+a}^{(k)}$ j = 5: Rate $a_{n+a}^{(k)}$		
	$j = 5$: Rate $a_{n+a}^{(k)}$		
	$j = 6$: Rate v_n		
xl(ndm,nen)	Element nodal reference coordinates		
ix(nen)	Element global node numbers		
tl(nen)	Element nodal temperature values		
s(nst,nst)	Element matrix (e.g., stiffness, mass)		
r(ndf,nen)	lf,nen) Element vector (e.g., residual, mass)		
	may also be used as $r(nst)$		
ndf	Number unknowns (max) per node		
ndm	Space dimension of mesh		
nst	Size of element arrays S and R		
	N.B. Normally $nst = ndf^*nen$		
isw	Task parameter to control computation		
	See prototype element in Figure 5.1		

Table 5.1: Arguments of FEAP Element Subprogram.

It is not necessary to implement all other tasks in an element, however, for those tasks that are not implemented it is important that the element routine not perform any calculations. Thus if the form of the branch is programmed as an IF-THEN-ELSE construct as shown in Fig. 5.1 then the ELSE should not carry out any operations *unless all options for ISW are programmed*. Similarly if the element is programmed using a SELECT-CASE form as

isw-Task	Type	Description	Access Command
0	Ο	Output label	SHOW,ELEM
1	R	Input d(*) parameters	Mesh:MATE,n
2	Ο	Check elements	Soln:CHECk
3	R	Compute tangent/residual	Soln:TANG
		Store in S/r	UTAN
4	Ο	Output element variables	Soln:STRE
5	Е	Compute cons/lump mass	Soln:MASS
		Store in S/r	MASS,LUMP
6	R	Compute residual	Soln:FORM,REAC
			Plot:REAC
7	Ο	Surface load/tangents	Mesh:SLOAd
8	Ο	Nodal projections	Soln:STRE,NODE
			Plot:STRE,PSTR
9	Ο	Damping	Soln:DAMP
10	Ο	Augmented Lagrangian update	Soln:AUGM
11	Ο	Error estimator	Soln:ERRO
12	R	History update	Soln:TIME
		For special history treatments else return	
13	Ο	Energy/momentum	Soln:TPLO,ENER
14	R	Initialize history	BATCh,INTEr
15	Ο	Body force	Mesh:BODY
16	Ο	J integrals	Soln: JINT
17	Ο	Set after activation	Soln:ACTI
18	Ο	Set after deactivation	Soln:DEAC
19		NOT AVAILABLE: used in modal/base	BASE
		Uses isw = 5 in element	
20	Ο	Element plotting	Plot:PELM
23	Ο	Compute element loads only	ARCL
25	Ο	Zienkiewicz-Zhu projection	Soln:ZZHU
26	R	Used to compute mesh boundary	Called by default.

Table 5.2: Task Options for FEAP Element Subprogram. R = Required; O = Optional; E = For eigensolutions

```
select case (isw)
    case(1)
c Input material parameters
    ...
    case default
    ...
end select
```

the CASE DEFAULT should not perform any operations unless all options are programmed. Finally, if the form

```
go to (1,2,.... ), isw
return
```

is used the RETURN statement should *always be included* as shown. This prevents any unexpected execution of a statement that appears after the GO TO.

Some of the options for additional data passed through common blocks is shown in Figure 5.2 with each variable defined in Table 5.3. Also, in Figure 5.3 the reference to common blocks using include statements is shown. In the prototype routine the number of nodes on an element (nen) which is used to dimension ul is passed in the labeled common /cdata/. Additional discussion is given below on use of some of the other data passed through the common blocks.

5.1 Material property storage

The material parameters to be stored in the array D with pointer np(25) may be input using the subprogram INPT2D. This subroutine is accessed by the statement:

CALL INPT2D(D,TDOF, NEV, TYPE)

where D is the array storing the material parameters; TDOF is returned as the parameter to access temperature; NEV is the number of element history variables to allocate to NH1; and TYPE is the element type. This routine inputs the commands as described in the user manual and stores the data for each material set into the D array elements as described in the following tables.

Variable	Definition
0	Page eject option
head	Title record
numnp	Number of mesh nodes
numel	Number of mesh elements
nummat	Number of material sets
nen	Maximum nodes/element
neq	Number active equations
ipr	Real variable precision
nstep	Total number of time steps
niter	Number of iterations current step
naugm	Number of augments current step
titer	Total iterations
taubm	Total augments
iaugm	Augmenting counter
iform	Number residuals in line search
dm	Element proportional load
n	Current element number
ma	Current element material set
mct	Print counter
iel	User element number
nel	Number nodes on current element
tt	Element stress values for TPLOt
bpr	Principal stretch
ctan	Element multipliers
ut	Element user values for TPLOt

Table 5.3: FEAP common block definitions.

5.2 Non-linear Transient Solution Forms

Before describing the steps in developing an element we summarize first the basic structure of the algorithms employed by *FEAP* to solve problems. Each problem to be solved using an ELMTnn routine is established in a standard finite element form as described in standard references (e.g., *The Finite Element Method*, 4th ed., by O.C. Zienkiewicz and R.L. Taylor, McGraw-Hill, London, 1989 (vol 1), 1991 (vol 2)). Here it is assumed this step leads to a set of non-linear ordinary differential equations expressed in terms of nodal displacements, velocities, and accelerations given by $\mathbf{u}_i(t)$, $\dot{\mathbf{u}}_i(t)$, and $\ddot{\mathbf{u}}_i(t)$, respectively. We denote the differential equation for node-*i* as the

Variable	Definition		
nh1	Pointer to t_n history data		
nh2	Pointer to t_{n+1} history data		
nh3	Pointer to element history		
ior	Current input logical unit		
iow	Current output logical unit		
nph	Pointer to global projection arrays		
ner	Pointer to global error indicator		
erav	Element error value		
j-int	J integral values		
ndf	Maximum dof/node		
ndm	Mesh space dimension		
nen1	Dimension 1 on IX array		
nst	Size of element matrix		
nneq	Total dof in problem		
ttim	Current time		
dt	Current time increment		
ci	Integration parameters		
hr	Real array data		
mr	Integer array data		

Table 5.4: FEAP common block definitions.

residual equation:

$$\mathbf{R}_i(\mathbf{u}_i(t), \dot{\mathbf{u}}_i(t), \ddot{\mathbf{u}}_i(t), t) = \mathbf{0}$$

To solve for the nodal displacements, velocities and accelerations it is necessary to introduce an algorithm to integrate the nodal quantities in time, specify a constitutive relation, and develop an algorithm to solve a (possibly) non-linear problem.

In *FEAP*, the integration method for nodal quantities is taken as a one step algorithm with each quantity defined only at discrete times t_n . Accordingly, we have displacements $\mathbf{u}_i(t_n)$ with velocities and accelerations denoted as

$$\dot{\mathbf{u}}_i(t_n) \approx \mathbf{v}_i(t_n)$$

and

$$\ddot{\mathbf{u}}_i(t_n) \approx \mathbf{a}_i(t_n)$$

A typical example for an integration algorithm for these discrete quantities is Newmark's method where

$$\mathbf{u}_{i}(t_{n+1}) = \mathbf{u}_{i}(t_{n}) + \Delta t \, \mathbf{v}_{i}(t_{n}) + \Delta t^{2} \left[\left(\frac{1}{2} - \beta\right) \mathbf{a}_{i}(t_{n}) + \beta \, \mathbf{a}_{i}(t_{n+1}) \right]$$

Parameter	Name	Description		
1	E	Young's modulus		
2	ν	Poisson ratio		
3	α	Thermal expansion coefficient		
4	ρ	Mass density		
5	-	Quadrature order for arrays		
6	-	Quadrature order for outputs		
7	a	Mass interpolation ($a = 0$: Diagonal; $a = 1$: Consistent		
8	q	Loading intensity (plates/shells)		
9	T_0	Stress free reference temperature		
10	κ	Shear factor (plates/shells/beams)		
11	b_1	Body force/volume in 1-directions		
12	b_2	Body force/volume in 2-directions		
13	b_3	Body force/volume in 3-directions		
14	h	Thickness (plates/shells)		
15	nh1	History variable counter		
16	stype	Two dimensional type: 1 - plane stress; 2 - plane strain;		
		$3 - axisymmetric^2$		
17	etype	Element formulation: 1 - displ; 2 - mixed; 3 - enhanced		
18	dtype	Deformation type: $\langle : \text{ finite}; \rangle$ small		
19	tdof	Thermal degree-of-freedom		
20	imat	Non-linear elastic material type		
21	d_{11}	Material moduli		
22	d_{22}	Material moduli		
23	d_{33}	Material moduli		
24	d_{12}	Material moduli		
25	d_{23}	Material moduli		
26	d_{31}	Material moduli		
27	g_{12}	Material moduli		
28	g_{23}	Material moduli		
29	g_{31}	Material moduli		
30	htype	Heat flag		

Table 5.5: Material Parameters

Parameter	Name	Description
31	ψ	Orthotropic angle x_1 principal axis 1
32	A	Area cross section (beam/truss)
33	I_{11}	Inertia cross section (beam/truss)
34	I_{22}	Inertia cross section (beam/truss)
35	I_{12}	Inertia cross section (beam/truss)
36	J	Polar inertia cross section (beam/truss)
37	κ_1	Shear factor plate
38	κ_2	Shear factor plate
39	-	Non-linear flag (beam/truss)
40	-	Inelastic material model type
41	Y_0	Initial yield stress (Mises)
42	Y_{∞}	Final yield stress (Mises)
43	β	Exponential hardening rate
44	H_{iso}	Isotropic hardening modulus (linear)
45	H_{kin}	Kinematic hardening modulus (linear)
46	-	Yield flag
47	β_1	Orthotropic thermal stress
48	β_2	Orthotropic thermal stress
49	β_3	Orthotropic thermal stress
50	-	Error estimator parameter
51	$ u_1 $	Viscoelastic shear parameter
52	$ au_1$	Viscoelastic relaxation time
53	ν_2	Viscoelastic shear parameter
54	$ au_2$	Viscoelastic relaxation time
55	$ u_3$	Viscoelastic shear parameter
56	$ au_3$	Viscoelastic relaxation time
57	nvis	Number of viscoelastic terms $(1-3)$
58	-	Damage limit
59	-	Damage rate
60	k	Penalty parameter

 Table 5.6:
 Material Parameters

Parameter	Name	Description	
61	K_1	Fourier thermal conductivity	
62	K_2	Fourier thermal conductivity	
63	K_3	Fourier thermal conductivity	
64	c	Fourier specific heat	
65	ω	Angular velocity	
66	Q	Body heat	
67	-	Heat constitution added indicator	
68	-	Follower loading indicator	
69	-	Rotational mass factor	
70	-	Damping factor	
71	g_1	Ground acceleration factor	
72	g_2	Ground acceleration factor	
73	g_3	Ground acceleration factor	
74	p_1	Ground acceleration proportional load number	
75	p_2	Ground acceleration proportional load number	
76	p_3	Ground acceleration proportional load number	
77	a_0	Rayleigh damping mass ratio	
78	a_1	Rayleigh damping stiffness ratio	
79	-	Plate/Shell/Rod shear activation flag	
80		Method: Type 1	
81		Method: Type 2	
82	-	Truss/Rod quadrature number	
83	-	Axial loading value	
84	-	Constitutive start indicator	
85	-	Polar angle indicator	
86	-	Polar angle coord_1	
87	-	Polar angle coord_2	
88	-	Polar angle coord_3	
89	-	Constitution transient type	
90	d_{31}	Plane stress recovery	
91	d_{32}	Plane stress recovery	
92	α_3	Plane stress recovery	

Table 5.7: Material Parameters

Parameter	Name	Description	
93	sref	Shear center type	
94	y_1	Shear center coordinate	
95	y_2	Shear center coordinate	
96	lref	Reference vector type	
97	n_1	Reference vector parameter	
98	n_2	Reference vector parameter	
99	n_3	Reference vector parameter	
100	-	Cross section shape type: $1 = \text{rectangles}; 2 = \text{tube};$	
		3 = Wide flange; $4 =$ Channel; $5 =$ Angle; $5 =$ Circle	
101-126	-	Shape data	
127	-	Surface convection (h)	
128	-	Free-stream temperature (T_{∞})	
129	-	Reference absolute temperature	
130	nseg	Number of hardening segments	
131-148	-	Segment data sets $e_p Y_{iso} H_{kin}$	
149	-	Total variables on frame section	
150	-	Piezoelectric flag	
151-159	-	Piezoelectric data	
160	-	Initial stress flag	
161-166	σ_{ij}	Initial stresses (constant)	
167	-	Tension/compression only indicator	
170	C	Fung pseudo elastic model modulus	
171	a_1	Fung model energy parameter	
172	a_2	Fung model energy parameter	
173	a_3	Fung model energy parameter	
174	a_4	Fung model energy parameter	
175	a_5	Fung model energy parameter	
176	a_6	Fung model energy parameter	
177	a_7	Fung model energy parameter	
178	a_8	Fung model energy parameter	
179	a_9	Fung model energy parameter	
180-181	-	Viscoplastic rate parameters	
182	-	Nodal quadrature parameters	
183	β_m	$\mathbf{M}_L - \mathbf{M}_C$ mass scaling factor	
184	c	Estimate on maximum wave speed	
185	-	Augmentation switch	
187		Implicit $= 0$; Explicit $= 1$ element integration	
188-200	-	Unused	

Table 5.8: Material Parameters

and

$$\mathbf{v}_{i}(t_{n+1}) = \mathbf{v}_{i}(t_{n}) + \Delta t \left[(1 - \gamma) \mathbf{a}_{i}(t_{n}) + \gamma \mathbf{a}_{i}(t_{n+1}) \right]$$

with \mathbf{u} , \mathbf{v} , and \mathbf{a} being the set of displacements, velocities, and accelerations at node-*i*, respectively.

A Newton method is commonly adopted to solve a non-linear (or linear) problem. To implement a Newton method it is necessary to linearize the residual equation. For FEAP, the Newton equation may be written as

$$\mathbf{R}_{i}^{(k+1)} = \mathbf{R}_{i}^{(k)} + \frac{\partial \mathbf{R}_{i}}{\partial \boldsymbol{\alpha}_{j}} \mid ^{(k)} d\boldsymbol{\alpha}_{j}^{(k)} = \mathbf{0}$$

where α_j is one of the variables at time t_{n+1} (e.g., $\mathbf{u}_j(t_{n+1})$). We define

$$\mathbf{S}_{ij}^{(k)} \;\;=\;\; - rac{\partial \mathbf{R}_i}{\partial oldsymbol lpha_j} \mid^{(k)}$$

and solve

$$\mathbf{S}_{ij}^{(k)} d \boldsymbol{\alpha}_j^{(k)} = \mathbf{R}_i^{(k)}$$

The solution is updated using

$${m lpha}_{j}^{(k+1)} \;\; = \;\; {m lpha}_{j}^{(k)} \; + \; d{m lpha}_{j}^{(k)} \; .$$

In the above (k) is the iteration number for the Newton algorithm. To start the solution for each step, FEAP sets

$$\boldsymbol{\alpha}_{j}^{(0)}(t_{n+1}) = \boldsymbol{\alpha}_{j}(t_{n})$$

where a quantity without the (k) superscript represents a converged value. For a linear problem, Newton's method converges in one iteration. Computing the residual after one iteration *must yield a zero value* to within the roundoff of the computer used. For non-linear problems, a properly implemented Newton's method *must exhibit a quadratic asymptotic rate of convergence*. Failure of the above performance for linear and non-linear cases implies a programming error in an implementation or lack of a consistently linearized algorithm (i.e., \mathbf{S}_{ij} is not an exact derivative of the residual).

In a non-linear problem, Newmark's method may be parameterized in terms of increments of displacement, velocity, or acceleration. From the Newmark formulas, the relations

 $d\mathbf{u}_i = \beta \,\Delta t^2 \, d\mathbf{a}_i$

and

$$d\mathbf{v}_i = \gamma \Delta t \, d\mathbf{a}_i$$

define the relationships between the increments. Note that only scalar multipliers involving β , γ , and Δt are involved between the different measures.

The tangent matrix for the transient problem using Newmark's method may be expressed in terms of the incremental displacement, velocity, or acceleration. As an example, consider the case where the solution is parameterized in terms of increments of the displacements (i.e., α_j is the displacement vector \mathbf{u}_j). For this case, the tangent matrix is (we do not show dependence on the iteration (k) for simplicity of notation)

$$\mathbf{S}_{ij} \, d\mathbf{u}_j = -\frac{\partial \mathbf{R}_i}{\partial \mathbf{u}_j} \, d\mathbf{u}_j - \frac{\partial \mathbf{R}_i}{\partial \mathbf{v}_k} \frac{\partial \mathbf{v}_k}{\partial \mathbf{u}_j} \, d\mathbf{u}_j - \frac{\partial \mathbf{R}_i}{\partial \mathbf{a}_k} \frac{\partial \mathbf{a}_k}{\partial \mathbf{u}_j} \, d\mathbf{u}_j \, .$$

Note that from the Newmark formulas

$$\frac{\partial \mathbf{a}_k}{\partial \mathbf{u}_j} = \frac{1}{\beta \, \Delta t^2} \, \boldsymbol{\delta}_{kj} \quad ; \quad \frac{\partial \mathbf{v}_k}{\partial \mathbf{u}_j} = \frac{\partial \mathbf{v}_k}{\partial \mathbf{a}_l} \frac{\partial \mathbf{a}_l}{\partial \mathbf{u}_j} = \frac{\gamma}{\beta \, \Delta t} \, \boldsymbol{\delta}_{kj}$$

in which δ_{kj} is the Kronnecker delta identity matrix for the k,j nodal pair . From the residual we observe that

$$\mathbf{K}_{ij} = -rac{\partial \mathbf{R}_i}{\partial \mathbf{u}_j} \quad ; \quad \mathbf{C}_{ij} = -rac{\partial \mathbf{R}_i}{\partial \mathbf{v}_j} \quad ; \quad \mathbf{M}_{ij} = -rac{\partial \mathbf{R}_i}{\partial \mathbf{a}_j}$$

define the tangent stiffness, damping, and mass, respectively. Thus, for the Newmark algorithm the total tangent matrix in terms of the incremental displacements is

$$\mathbf{S}_{ij} = \mathbf{K}_{ij} + rac{\gamma}{\beta \Delta t} \mathbf{C}_{ij} + rac{1}{\beta \Delta t^2} \mathbf{M}_{ij} .$$

For other choices of increments, the tangent may be written in the general form

$$\mathbf{S}_{ij} = c_1 \, \mathbf{K}_{ij} + c_2 \, \mathbf{C}_{ij} + c_3 \, \mathbf{M}_{ij}$$

where the c_i are scalar quantities involving the integration parameters of the method selected and Δt . Thus, any one step integrator may be considered and will affect only the specification of the constants in the tangent.

In *FEAP* the element tangent matrix, S_{ij} , is stored as a two dimensional array which is dimensioned as s(nst,nst), where nst is the product of ndf and nen, with ndf the maximum number of degree-of-freedoms at any node in the problem and nen the maximum number of nodes on any element. The ordering of the unknowns into nst must be carefully aligned in order for FEAP to properly assemble each element matrix into the global tangent. The ordering is such that sub-matrices are defined for each node attached to the element. Thus

$$\mathbf{S} = \begin{bmatrix} \mathbf{S}_{11} & \mathbf{S}_{12} & \mathbf{S}_{13} & \cdots \\ \mathbf{S}_{21} & \mathbf{S}_{22} & \mathbf{S}_{23} & \cdots \\ \mathbf{S}_{31} & \mathbf{S}_{32} & \mathbf{S}_{33} & \cdots \\ \cdots & \cdots & \cdots & \cdots \end{bmatrix}$$

where \mathbf{S}_{ij} is the sub-matrix for nodal pairs i, j. Each of the sub-matrices is a square matrix of the size of the maximum number of degree-of-freedoms in the problem which is passed to the subprogram as ndf. Thus,

$$\mathbf{S}_{ij} = \begin{bmatrix} S_{11}^{ij} & S_{12}^{ij} & S_{13}^{ij} & \cdots \\ S_{21}^{ij} & S_{22}^{ij} & S_{23}^{ij} & \cdots \\ S_{31}^{ij} & S_{32}^{ij} & S_{33}^{ij} & \cdots \\ \cdots & \cdots & S_{ndf,ndf}^{ij} \end{bmatrix}$$

in which S_{ab}^{ij} is an array coefficient for nodal pair i, j for the degree-of-freedom pair a, b.

In FEAP, the element residual may be stored as one dimensional array which is dimensioned r(nst) with entries stored in the same order as the rows of the element tangent matrix or as a two dimensional array which is dimensioned as r(ndf,nen). The one dimensional form of the residual is given as

$$\mathbf{R} = egin{bmatrix} \mathbf{R}_1 \ \mathbf{R}_2 \ \mathbf{R}_3 \ dots \end{bmatrix}$$

where the entries in each submatrix are given as

$$\mathbf{R}_{i} = \begin{bmatrix} R_{1}^{i} \\ R_{2}^{i} \\ R_{3}^{i} \\ \vdots \\ R_{ndf}^{i} \end{bmatrix}$$

The two dimensional form places the entries \mathbf{R}_i as columns. Accordingly,

$$\mathbf{R}~=~\begin{bmatrix} \mathbf{R}_1 & \mathbf{R}_2 & \mathbf{R}_3 & \cdots \end{bmatrix}$$
 .

The two forms for defining the residual \mathbf{r} are equivalent based on the Fortran ordering of information into double subscript arrays.

If ndf is larger than needed for the element and residual the unused positions need not be defined (the tangent array s and the residual r are set to zero before each element routine is called).

The arrays xl(i,j), ul(i,j,1), ul(i,j,4) and ul(i,j,5) (described in Table 5.1) are used to obtain the nodal coordinates, displacements, velocities and accelerations, respectively. When *FEAP* solves a problem without transient loading (e.g., inertial

Parameter	Description		
$\operatorname{ctan}(1)$	c_1 : Multiplier of s matrix for ul(i,j,1) terms		
	(e.g., stiffness matrix multiplier)		
$\operatorname{ctan}(2)$	c_2 : Multiplier of s matrix for ul(i,j,4) terms		
	(e.g., damping matrix multiplier)		
$\operatorname{ctan}(3)$	c_3 : Multiplier of s matrix for ul(i,j,5) terms		
	(e.g., mass matrix multiplier)		

 Table 5.9:
 Tangent Parameters

loading as mass times acceleration) the velocities and accelerations are set to zero prior to calling the element subroutine. Consequently, in programming the steps to compute the residual \mathbf{r} the inertia terms have no effect for static or quasi-static problems and may be included (generally there are very few additional operations involved to add these terms). The programming of the tangent array, however, must distinguish between cases in which transient (e.g., inertial) loads are present and those in which they are omitted. The different cases are implemented in *FEAP* by making appropriate assignments to the c_i parameters. To facilitate the programming of the tangent array returned in \mathbf{s} for the various cases, a parameter array ctan(3) is passed to the subprogram in labeled common eltran. When the task parameter isw is 3, the values in the ctan array are interpreted according to Table 5.9.

Thus, in solid mechanics applications the tangent matrix is defined in an element routine as

$$\mathbf{S} = ctan(1) \mathbf{K} + ctan(2) \mathbf{C} + ctan(3) \mathbf{M}$$

where \mathbf{K} is the stiffness matrix, \mathbf{C} is the damping matrix, and \mathbf{M} is the mass matrix. For non-linear applications these matrices normally are computed with respect to the current values of the available solution parameters. The values provided in the ctan array are set by *FEAP* according to the active transient solution option. For a static option both ctan(2) and ctan(3) are zero. For options integrating first order differential equations in time only ctan(3) will be zero. For options integrating second order differential equations in time all the parameters are non-zero.

5.3 Example: 2-Node Truss Element

An element routine carries out tasks according to the value assigned to the parameter isw as indicated in Table 5.2 To describe basic steps to program the various tasks

defined by isw, we consider next the problem of a 2-node, linear elastic truss element for small deformation applications. The element is described in sufficient generality to permit solution of both two and three dimensional truss problems.

5.3.1 Theory for a Truss

The governing equations for a typical truss member element, shown in Figure 5.4, are the balance of momentum equation:

$$\frac{\partial (A\sigma_{ss})}{\partial s} + A b_s = \rho A \ddot{u}_s$$

the strain-displacement equation for small deformations:

$$\epsilon_{ss} = \frac{\partial u_s}{\partial s}$$

and a constitutive equation. For example, considering a linear elastic material the constitutive equation may be written as

$$\sigma_{ss} = E \,\epsilon_{ss}$$

Boundary and initial conditions must also be specified to obtain a well posed problem; however, our emphasis here is the derivation of the element arrays associated with the above differential equations. In the above:

- s is the coordinate along the truss member axis,
- b_s is a loading in direction s per unit length,
- A is the truss cross-section area,
- ρ is the mass density per unit volume,
- u_s is a displacement in direction s,
- \dot{v}_s is an acceleration in direction $s \ (v = \dot{u}),$
- ϵ_{ss} is a strain along the truss member axis, and
- σ_{ss} is the stress on a truss cross section.

The equations may also be deduced from the variational equation

$$\delta\Pi = \int_L \delta\epsilon_{ss} \sigma_{ss} A \, ds + \sum_{i=1}^d \int_L \delta u_i \rho A \, \dot{v}_i \, ds - \sum_{i=1}^d \int_L \delta u_i \, b_i \, ds + \delta \Pi_{ext}$$

where $\delta \Pi_{ext}$ contains the boundary and loading terms not associated with an element. Where, in addition to previously defined quantities, we define:

- d is the spatial dimension of the truss (1, 2, or 3),
- x_i are the Cartesian coordinates in the *d* directions.
- L is the length of the truss member,
- δu_i is a virtual displacement in direction x_i ,
- \dot{v}_i is an acceleration in direction x_i $(v = \dot{u})$,
- b_i is a loading in direction x_i per unit length, and
- $\delta \epsilon_{ss}$ is a virtual strain along the truss axis.

For a straight truss member the displacement along the axis, u_s may be expressed in terms of the components in the directions x_i as

$$u_s = \mathbf{l} \cdot \mathbf{u}(s, t) = \sum_{i=1}^d l_i u_i(s, t)$$

where t is time, **u** is the displacement vector with components u_i , **l** is a unit vector along the axis of the member with direction cosines l_i defined by

$$l_i = \frac{\partial x_i}{\partial s} = \frac{x_{i2} - x_{i1}}{L}$$
$$L^2 = \sum_{i=1}^d (x_{i2} - x_{i1})^2$$

and x_{i1} , x_{i2} are the coordinates of nodes 1 and 2, respectively. The displacement components are interpolated on the 2-node truss member as

$$u_i(s, t) = (1 - \xi) u_{i1}(t) + \xi u_{i2}(t) ; \xi = \frac{s}{L}$$

in which u_{i1} , u_{i2} are the displacements at nodes 1 and 2. The virtual displacements are obtained from the above by replacing u_i by δu_i , etc. The truss strain is

$$\epsilon_{ss} = \frac{\partial u_s}{\partial s} = \sum_{i=1}^d \, l_i \, \frac{\partial u_i}{\partial s} \, .$$

Using the interpolations for the displacement components yields

$$\epsilon_{ss} = \frac{1}{L^2} \sum_{i=1}^d \Delta x_i \, \Delta u_i$$

where

$$\Delta x_i = x_{i2} - x_{i1} = l_i L$$

and

$$\Delta u_i = u_{i2} - u_{i1} \; .$$

Thus, in matrix form the strain is

$$\epsilon_{ss} = \frac{1}{L^2} \sum_{i=1}^{d} \begin{bmatrix} -\Delta x_i & \Delta x_i \end{bmatrix} \begin{bmatrix} u_{i1} \\ u_{i2} \end{bmatrix}$$

Using the above displacement interpolations, the variational equation for the truss may be expressed in matrix form as

$$\begin{split} \delta\Pi &= \begin{bmatrix} \delta u_{i1} & \delta u_{i2} \end{bmatrix} \left\{ \int_L \frac{1}{L^2} \begin{bmatrix} -\Delta x_i \\ \Delta x_i \end{bmatrix} \sigma_{ss} A ds + \int_L \begin{bmatrix} 1-\xi \\ \xi \end{bmatrix} \rho A \begin{bmatrix} 1-\xi \\ \xi \end{bmatrix} ds \begin{bmatrix} \ddot{u}_{i1} \\ \ddot{u}_{i2} \end{bmatrix} \\ &- \int_L \begin{bmatrix} 1-\xi \\ \xi \end{bmatrix} b_i ds \right\} + \delta\Pi_{ext} \; . \end{split}$$

FEAP constructs the finite element arrays from the element residuals which are obtained from the negative of the terms multiplying the nodal displacements. Accordingly,

$$\mathbf{R}_{i} = \begin{bmatrix} R_{i1} \\ R_{i2} \end{bmatrix} = \int_{L} \begin{bmatrix} 1 - \xi \\ \xi \end{bmatrix} b_{i} \, ds$$
$$- \int_{L} \frac{1}{L^{2}} \begin{bmatrix} -\Delta x_{i} \\ \Delta x_{i} \end{bmatrix} \sigma_{ss} \, A \, ds - \int_{L} \begin{bmatrix} 1 - \xi \\ \xi \end{bmatrix} \rho \, A \begin{bmatrix} 1 - \xi \\ \xi \end{bmatrix} \, ds \begin{bmatrix} \ddot{u}_{i1} \\ \ddot{u}_{i2} \end{bmatrix}$$

is the residual for the i-coordinate direction. For constant properties and loading over an element length (note that for this case the stress will also be constant since strains are constant on the element), the above may be integrated to yield

$$\mathbf{R}_{i} = \begin{bmatrix} R_{i1} \\ R_{i2} \end{bmatrix} = \frac{1}{2} b_{i} L \begin{bmatrix} 1 \\ 1 \end{bmatrix} - \frac{\sigma_{ss} A}{L} \begin{bmatrix} -\Delta x_{i} \\ \Delta x_{i} \end{bmatrix} - \frac{\rho A L}{6} \begin{bmatrix} 2 & 1 \\ 1 & 2 \end{bmatrix} \begin{bmatrix} \ddot{u}_{i1} \\ \ddot{u}_{i2} \end{bmatrix} .$$
(5.1)

For the present we assume the material model is a linear elastic in which the stress is related to strain through

$$\sigma_{ss} = E \,\epsilon_{ss}$$

where E is the Young's modulus.

Based on a linear elastic material, the term in the residual involving σ_{ss} may be written as

$$\frac{\sigma_{ss}A}{L} \begin{bmatrix} -\Delta x_i \\ \Delta x_i \end{bmatrix} = \frac{EA}{L^3} \begin{bmatrix} -\Delta x_i \\ \Delta x_i \end{bmatrix} \sum_{j=1}^d \begin{bmatrix} -\Delta x_j & \Delta x_j \end{bmatrix} \begin{bmatrix} u_{j1} \\ u_{j2} \end{bmatrix} .$$

For the linear elastic material, a stiffness matrix may be expressed as

$$\mathbf{K}_{ij} = \frac{EA}{L^3} \begin{bmatrix} -\Delta x_i \\ \Delta x_i \end{bmatrix} \begin{bmatrix} -\Delta x_j & \Delta x_j \end{bmatrix} = \begin{bmatrix} k_{ij} & -k_{ij} \\ -k_{ij} & k_{ij} \end{bmatrix}$$

where

$$k_{ij} = \frac{EA}{L^3} \Delta x_i \Delta x_j \; .$$

The residual may now be written using a stiffness and mass matrix as

$$\mathbf{R}_{i} = \begin{bmatrix} R_{i1} \\ R_{i2} \end{bmatrix} = \frac{1}{2} b_{i} L \begin{bmatrix} 1 \\ 1 \end{bmatrix} - \sum_{j=1}^{d} \begin{bmatrix} k_{ij} & -k_{ij} \\ -k_{ij} & k_{ij} \end{bmatrix} \begin{bmatrix} u_{j1} \\ u_{j2} \end{bmatrix} - \begin{bmatrix} m_{11} & m_{12} \\ m_{21} & m_{22} \end{bmatrix} \begin{bmatrix} \ddot{u}_{i1} \\ \ddot{u}_{i2} \end{bmatrix}$$
(5.2)

with

$$m_{11} = m_{22} = \frac{\rho A L}{3}$$
; $m_{12} = m_{21} = \frac{\rho A L}{6}$

For non-linear material behavior the residual must be computed using Equation 5.1 with the stress replaced by the value computed from the constitutive equation.

The integration method for nodal quantities is taken as Newmark's method described in Section 5.2. The residual and tangent matrix for a Newton type method are now available and may be inserted into \mathbf{R} and \mathbf{S} after noting that for the truss that the damping matrix \mathbf{C} is zero. The residual may be programmed directly from Equation 5.1 and an implementation using the two dimensional form r(ndf,nen) is shown in Figure 5.5.

Similarly, using the results from Section 5.2, the tangent matrix for the truss may be programmed as indicated in Figures 5.6 and 5.7.

5.4 Additional Options in Elements

FEAP permits some additional options to be included within element tasks.

5.4.1 Task 1 Options

Often it is necessary to use several element types to perform an analysis. For example it may be necessary to use both truss and frame (bending resistant) elements to perform an analysis. As developed in Section 5.3, the truss element has one degree of freedom for each spatial dimension, whereas, the frame element must have additional unknowns to represent the bending behavior. For nodes connected only to truss elements it is not necessary to have the additional degrees-of-freedom active and a user would be required to specify restraint conditions for these nodes and degrees-of-freedom. By inserting the following lines of code into the truss element routine for the isw = 1 task *FEAP* will automatically eliminate any unneeded degrees-of-freedom.

```
do i = ndm+1,ndf
ix(i) = 0
end do ! i
```

Note that for isw = 1 the ix parameter is not used to pass the nodal connection array but is used to return the list of unused degrees-of-freedom.

Utility routines are also provided to assist users in providing the necessary list of nodes needed to properly draw the mesh each element type during plot outputs. The names of the routines are listed in Table 5.10 and each routine is called as

```
call plname (iel)
```

where iel is an integer parameter defined in common eldata. If no call to a subprogram is included each element is assumed to be a 4 to 9 node quadrilateral and default drawing order will be assigned. Users may construct their own drawing order also by following the steps employed in one of the routines defined in Table 5.10.

5.4.2 Task 6 Options

The TPLOt solution command includes an option to save specific element quantities (e.g., stress, strain, etc.). This option is implemented for user elements by including the common

Routine Name	Description
PLTLN2	2-node line element
PLTRI3	3-node triangular element
PLQUD4	4-node quadrilateral element
PLTRI6	6-node triangular element
PLTET4	4-node tetrahedron element
PLBRK8	8-node brick element

Table 5.10: Element Plot Definition Subprograms

real*8 tt common /elplot/ tt(100)

and then inserting the statement

tt(i) = value

at an appropriate location in the isw = 3 task.

For example if it were desired to save the force and strain in the truss element the statements

tt(1)	=	EA*eps	!	Element	axial	force
tt(2)	=	eps	!	Element	axial	strain

could be placed anywhere after the stress and strain are defined. These values would be output by using a solution command sequence such as

```
batch
  tplot
end
stress,nn,1 ! saves force for element nn
stress,nn,2 ! saves strain for element nn
show ! writes tplot items to output file
```

5.5 Projection of element variables to nodes

The STREss, NODE solution command and the PLOT, STRE command require a projection of element variables to nodes. A continuous stress field is assumed to obtain the nodal

values. Accordingly,

$$\sigma = N_{\alpha} \, \sigma_{\alpha}$$

where σ is any value which is to be projected to nodes (e.g., a stress or strain), N_{α} are shape functions for the element type considered, and σ_{α} nodal values of the projected quantity. The projection routine uses a diagonal weight matrix to project the values. For simple elements the matrix is computed by a procedure identical to mass lumping. For example,

$$M_{\alpha\alpha} = \int_{\Omega} N_{\alpha} \,\mathrm{d}\Omega$$

defines a 'row sum' form of projection^[2, 3]. Using the above results in the set of equations and a least square fit with the finite element values $\hat{\sigma}$ gives the equation set

$$M_{\alpha\alpha}\,\sigma_\alpha = \int_\Omega \,N_\alpha\,\hat\sigma\,\mathrm{d}\Omega~.$$

This defines nodal values for projected quantities. Since the coefficient matrix is diagonal the solution to the set of equations for each component is trivial. The actual solution is performed automatically by *FEAP*.

To permit each element to project its own quantities it is necessary to add the projection operations for each element under $ISW = 8.^3$ These are performed locally for each element similar to all other operations. Figure 5.8 shows a simple routine for two-dimensional elements with 4-stress components begin projected. When multiple element types are used in an analysis users must be careful to project like quantities to common values of the ST(nen,*) array so as to get correct results. Also, when results are displayed it is necessary to plot results by material type to obtain correct indications of stress discontinuities at material interfaces.

5.6 Elements with History Variables

FEAP provides options for each element to manage variables which must be saved during the solution. These are history variables and are separated into three groups: (a) Variables associated with the last converged solution time t_n ; (b) Variables associated with the current solution time t_{n+1} ; and variables which are not associated to any particular time. All history variables are associated with the allocation name H which has a pointer value 49. Users are not permitted direct access to the data stored as H (of course, it is possible to access from hr(np(49)) but this should not normally

³An implementation of the Zienkiewicz-Zhu projection method is implemented using ISW = 24.

be attempted!). Before calling the element routine for each element, *FEAP* transfers the required history variable to a local storage for each type. Users may then access the history data for each element and if necessary update values and return them *FEAP*. Only for specific actions will the local history data be transferred back to the appropriate H locations. The element history data associated with t_n starts at the memory address of the pointer for NH1 using the double precision dummy array HR in blank common; similarly data for t_{n+1} starts at the memory address of the pointer for NH2, and that not associated with a time at NH3. The three pointers are passed to each element routine in the labeled common

```
integer nh1,nh2,nh3
common /hdata/ nh1,nh2,nh3
```

5.6.1 Assigning amount of storage for each element

The specification for the amount of history information to be associated with each material set is controlled in the isw = 1 task of an element routine. For each material type specified within the element routine a value for the length of the NH1 and the NH3 data must be provided (the amount of NH2 data will be the same as for NH1). This is accomplished by setting the variables nh1 and nh2 in common hdata (see above) to the required values. That is, the statements required are:

if(isw .eq. 1) then
 . . .
 nh1 = 6
 nh3 = 10
 . . .

reserves 6 words of NH1 and NH2 data and 10 words of NH3 data for each element with the current material number. Care should be taken to minimize the number of history variables since, for very large problems, the memory requirements can become large, thus reducing the size of problem that *FEAP* can solve.

5.6.2 Accessing history data for each element

As noted above the data for each element is contained in arrays whose first word is located at hr(nh1), hr(nh2) (where nh1 and nh2 are pointers) for t_n , t_{n+1} , respectively; and at hr(nh3) for that not associated with time (note that there are values for each only if non-zero values are assigned to nh1 and/or nh3 during the isw = 1 task. Any other allocated data follows immediately after each first word It is a users responsibility to manage what is retained in each variable type; however, the order of placing the t_n and t_{n+1} data into the NH1 and NH2 arrays should be identical. There are no provisions to store integer history variables separately from double precision quantities. It is necessary to cast the integer data as double precision and move to the history location. For example, using the statement

hr(nh3+5) = dble(ivarbl)

saves the value for the integer variable ivarbl in the sixth word of the NH3 element history array. At a subsequent iteration for this element the value of the integer would be recovered as

ivarbl = int(hr(nh3+5))

While this wastes storage for integer variables, experience indicates there is little need to save many integer quantities and, thus, it was not deemed necessary to provide for integer history variables separately.

Although users may define new values for any of the hr(nh1), hr(nh2), or hr(nh3) types, the new quantities will be returned to the H history for the element only for isw tasks where residuals are being formed for a solution step (i.e., solution command FORM, TANG, 1, or UTAN, 1 and for history reinitialization during a time update (i.e., solution command, TIME). These access the task options isw equal to 3 or 6 and 14, respectively.

If a user adds a new option for which it is desired to save the history variables, it is necessary to set the variables hflgu and h3flgu to true as required, if no update is wanted the variables should be set to false. These parameters are located in

logical	L	hflgu,h3fl	.gu
$\verb common $	/hdatam/	hflgu,h3fl	gu

5.7 Elements with Finite Rotation Parameters

When considering structural elements that undergo large displacements it is usually necessary to treat the rotation parameters for large angle changes. The nodal parameters for this case are a rotation vector $\boldsymbol{\theta}$ and the finite rotations are given as an

orthogonal matrix Λ .

$$\Lambda_{n+1} = \exp[\hat{\theta}] \Lambda_n$$

in which $\hat{\boldsymbol{\theta}}$ denotes a skew matrix given as

$$\hat{\boldsymbol{\theta}} = \begin{bmatrix} 0 & \theta_3 & -\theta_2 \\ -\theta_3 & 0 & \theta_1 \\ \theta_2 & -\theta_1 & 0 \end{bmatrix}$$

The actual method used to update the rotations and their increments must be specified when writing the element module ELMTnn and is performed by a user subprogram named UROTmm where mm is a number between 01 and 10. To specify which routine is to be used, it is necessary to include the statement

rotyp = mm

in the section of ELMTnn where isw = 1. This parameter is located in the common erotas which has the structure:

	real*8	xln				
	real*8	rots	,rvel	,racc	,thkl	
	integer					rotyp
	<pre>common /erotas/</pre>	xln(9,9,4),				
,	&	rots(3,9,2)	,rvel(3,9,2)	,racc(3,9,2)	,thkl(9)	,rotyp

The other entries in the common are arrays that return values for each element to treat the rotation values and rates. We shall return to their description after describing the treatment of the global nodal data for rotations.

5.7.1 Nodal rotation treatment: UROTmm subprogram

The nodal rotation data is stored in the array xlg which is dimensioned as

xlg(9,6,numnp)

For node ng, the entries in xlg are stored as follows:

Component	I/O	Description
XLG(1:9,1,ng)	Ι	Rotation matrix $\mathbf{\Lambda}_n$ at time t_n
XLG(1:9,2,ng)	0	Rotation matrix $\mathbf{\Lambda}_{n_a}$ at time t_{n_a}
XLG(1:9,3,ng)	0	Rotation matrix $\mathbf{\Lambda}_{n+1}$ at time t_{n+1}
XLG(1:3,4,ng)	0	Rotation increment angle $\Delta \boldsymbol{\theta}$
XLG(4:6,4,ng)	Ι	Rotation rate $\boldsymbol{\omega}_n$ at time t_n
XLG(7:9,4,ng)	Ι	Rotation acceleration $\boldsymbol{\alpha}_n$ at time t_n
XLG(1:3,5,ng)	0	Rotation angle $\boldsymbol{\theta}$
XLG(4:6,5,ng)	0	Rotation acceleration $\boldsymbol{\alpha}_{n+1}$ at time t_{n+1}
XLG(7:9,5,ng)	0	Rotation acceleration $\boldsymbol{\alpha}_{n+a}$ at time t_{n+a}
XLG(1:9,6,ng)	-	Rotation matrix $\mathbf{\Lambda}_0$ at time t_0

While storage is provided for the 3×3 rotation matrices the representation may also be specified in terms of quaternions for which only 4 components are necessary. In this case the 9 entries may be divided into two 4 entry quantities if required. Indeed, the space may be used in anyway necessary provided, no conflict in the way each time value is associated to the data. Note that sufficient storage is available to define integration methods for which the rotation is defined at an intermediate time t_{n+a} .

For a typical node ${\tt n}$ in the mesh the location of the entries in the ${\tt xlg}$ array are obtained from

ng = mropt(n, 2)

and the routine UROTmm is called as:

call urotmm(xlg(1,1,ng),xlg(1,2,ng),xlg(1,3,ng),
& xlg(1,4,ng),xlg(1,5,ng),
& xlg(4,4,ng),xlg(4,5,ng),
& xlg(7,4,ng),xlg(7,5,ng),du(j,n),tsw)

where du(j,n) are the solution increments from the solver and tsw is the time update switch which is set according to

tsw = 1: Initialize for new time step tsw = 2: Update within a time step tsw = 3: Back up to beginning of time step

The entry u(j,n) is the location for the first entry in the rotation vector $\boldsymbol{\theta}$.

5.7.2 Local nodal rotation treatment

When each element that is associated with nodal rotation parameters is called the rotation data is transferred to local storage in a manner similar to treatment of translations. The local data is passed to each element using the common **erotas** defined above. The entries in the local arrays are extracted from the global array according to:

xln(1:9,nl,1)	= xlg(1:9,1,ng)
xln(1:9,nl,2)	= xlg(1:9,2,ng)
xln(1:9,nl,3)	= xlg(1:9,3,ng)
xln(1:9,nl,4)	= xlg(1:9,6,ng)
rots(1:3,nl,1)	= xlg(1:3,4,ng)
rots(1:3,nl,2)	= xlg(1:3,5,ng)
rvel(1:3,nl,1)	= xlg(4:6,4,ng)
rvel(1:3,nl,2)	= xlg(4:6,5,ng)
<pre>racc(1:3,nl,1)</pre>	= xlg(7:9,4,ng)
racc(1:3,nl,2)	= xlg(7:9,5,ng)

where **nl** is a local node number between 1 and 9 (the maximum provided in the current **erotas** and **ng** is the global node number associated with each local number.

Using the above data structure one can program the updates in any manner that does not conflict with the time treatment. The only interface to FEAP is through how the increment du(4:5,n) is defined.

5.8 Energy Computation

FEAP elements provide an option to accumulate the total momenta and energy during the solution process. The values are accumulated in the array EPL(20) when the switch parameter isw is 13 and written to a file named Pxxxx.ene (where xxxx is extracted from the problem input filename) whenever the solution command TIME is used. The array EPL(2) is in the common block named ptdat6 which has the structure:

real*8 epl integer iepl, neplts common /ptdat6/ epl(20),iepl(2,20),neplts

For problems in solid mechanics the linear momenta are stored as follows:

Component	Description
EPL(1) - EPL(3)	Linear momenta
EPL(4) - EPL(6)	Angular momenta
EPL(7)	Kinetic energy
EPL(8)	Stored energy
EPL(9)	Work by external loads
EPL(10)	Total energy

Table 5.11: Momenta and Energy Assignments

The linear momenta are computed as:

$$\mathbf{p} = \int_{\Omega} \rho \mathbf{v} \, d\Omega$$

the angular momenta as:

$$\boldsymbol{\pi} = \int_{\Omega} \left(\mathbf{I} \, \boldsymbol{\omega} + \mathbf{x} \times \mathbf{p} \right) d\Omega$$

the kinetic energy

$$K = \int_{\Omega} \rho \mathbf{v} \cdot \mathbf{v} \ d\Omega$$

the stored energy as

$$U = \int_{\Omega} W(\mathbf{C}) \ d\Omega$$

and the work by external loads as

$$V = \int_{\Gamma} (\mathbf{x} - \mathbf{X}) \cdot \mathbf{F}_{ext} \ d\Gamma \ .$$

The value of the displacement and velocity at the current time t_{n+1} are passed in ul(i,j,1) and ul(i,j,4), respectively. Note that this is true no matter which time integration algorithm is specified.

5.9 A Non-linear Theory for a Truss

A simple non-linear theory for a two or three dimensional truss which may undergo large displacements for which the strains remain small may be developed by defining the axial strain approximation in each member as

$$\epsilon_{ss} = \frac{\partial u_s}{\partial s} + \frac{1}{2} \sum_{j=1}^{d-1} \left(\frac{\partial u_{nj}}{\partial s} \right)^2$$

where u_{nj} is a displacement component normal to the axis of the member. The virtual strain from a linearization of the strain is given as

$$\delta \epsilon_{ss} = \frac{\partial \delta u_s}{\partial s} + \sum_{j=1}^{d-1} \left(\frac{\partial \delta u_{nj}}{\partial s} \right) \left(\frac{\partial u_{nj}}{\partial s} \right) \,.$$

An algorithm to define the two orthogonal unit vectors which are normal to the member may be constructed by taking

$$\mathbf{v} = \mathbf{e}_k$$

where k is a direction for which a minimum value of the direction cosine l_i exists (for a 2-dimensional problem defined in the x_1 , x_2 plane **v** may be taken as \mathbf{e}_3). Now,

$$\mathbf{n}_1 \;=\; rac{\mathbf{v} imes \mathbf{l}}{\mid \mathbf{v} imes \mathbf{l} \mid}$$

and

$$\mathbf{n}_2 = \mathbf{l} \times \mathbf{n}_1$$
 .

Using these vectors the two normal components of the displacement are given by

$$u_{nj}(s, t) = \mathbf{n}_j \cdot \mathbf{u}(s, t) = \sum_{i=1}^d n_{ji} u_i(s, t)$$

and the derivative by

$$\frac{\partial u_{nj}}{\partial s} = \sum_{i=1}^d n_{ji} \frac{\partial u_i}{\partial s} .$$

Collecting terms and combining with previously defined quantities the virtual strain may be written as

$$\delta \epsilon_{ss} = \frac{\partial \delta \mathbf{u}}{\partial s} \cdot \left[\mathbf{g} \right]$$

where

$$\mathbf{g} = \mathbf{l} + \sum_{j=1}^{d-1} \frac{\partial u_{nj}}{\partial s} \mathbf{n}_j$$

After differentiation of the displacement field the discrete form of the virtual strain is given by

$$\delta \epsilon_{ss} = rac{1}{L} \begin{bmatrix} \delta \mathbf{u}_1 & \delta \mathbf{u}_2 \end{bmatrix} \cdot \begin{bmatrix} - \mathbf{g} \\ \mathbf{g} \end{bmatrix} .$$

Substituting the above virtual strain expression into the weak form gives the modified residual expression

$$\mathbf{R}_{i} = \frac{1}{2} b_{i} L \begin{bmatrix} 1\\1 \end{bmatrix} - \sigma_{ss} A \begin{bmatrix} -g_{i}\\g_{i} \end{bmatrix} - \rho A \frac{L}{6} \begin{bmatrix} 2 & 1\\1 & 2 \end{bmatrix} \begin{bmatrix} \ddot{u}_{i1}\\\ddot{u}_{i2} \end{bmatrix} .$$
(5.3)

The tangent tensor is obtained by linearizing the residual as shown previously. The only part which is different is the term with σ_{ss} . Noting that

$$d\epsilon_{ss} = \left[\mathbf{g} \right] \cdot \frac{\partial d\mathbf{u}}{\partial s}$$

and

$$d\,\delta\epsilon_{ss} = \frac{\partial\delta\mathbf{u}}{\partial s}\,\cdot\,(\mathbf{n}_1\otimes\mathbf{n}_1\ +\ \mathbf{n}_2\otimes\mathbf{n}_2)\,\cdot\,\frac{\partial d\mathbf{u}}{\partial s}$$

If the \mathbf{n}_i are constructed as *column* vectors then the tensor product becomes a matrix defined as

$$\mathbf{G} = \mathbf{n}_1 \otimes \mathbf{n}_1 + \mathbf{n}_2 \otimes \mathbf{n}_2 = \mathbf{n}_1 \mathbf{n}_1^T + \mathbf{n}_2 \mathbf{n}_2^T$$

With these definitions, the *tangent* matrix for the non-linear problem is given as

$$\mathbf{K}_{ij} = \frac{EA}{L} \begin{bmatrix} -g_i \\ g_i \end{bmatrix} \begin{bmatrix} -g_j & g_j \end{bmatrix} + \frac{\sigma_{ss}A}{L^2} \begin{bmatrix} G_{ij} & -G_{ij} \\ -G_{ij} & G_{ij} \end{bmatrix}$$

Notice that for the linear problem

$$g_i = \frac{\Delta x_i}{L}$$

thus, the only difference between the linear and non-linear problem is the definition of ϵ_{ss} in terms of displacements, the modification for geometric effects for the g_i and the second term on the tangent matrix which is sometimes called the *geometric* stiffness part.

character*4 o,head common /bdata/ o,head(20) integer numnp,numel,nummat,nen,neq,ipr common /cdata/ numnp,numel,nummat,nen,neq,ipr integer nstep,niter,naugm, titer,taugm,tform common /counts/ nstep,niter,naugm, titer,taugm,tform iaugm, iform, intvc, iautl, nstepa integer common /counts/ iaugm, iform, intvc, iautl, nstepa real*8 dm n,ma,mct,iel,nel,pstyp integer common /eldata/ dm,n,ma,mct,iel,nel,pstyp real*8 tt common /elplot/ tt(200) real*8 bpr, ctan, psil common /eltran/ bpr(3),ctan(3),psil real*8 ut common /eluser/ ut(200) nh1,nh2,nh3,ht1,ht2,ht3, nlm,plm integer common /hdata/ nh1,nh2,nh3,ht1,ht2,ht3, nlm,plm ior, iow, ilg integer common /iofile/ ior,iow,ilg integer nph,ner ,jshft real*8 erav,j_int common /prstrs/ nph,ner,erav,j_int(3),jshft integer ndf,ndm,nen1,nst,nneq,ndl,nnlm,nadd common /sdata/ ndf,ndm,nen1,nst,nneq,ndl,nnlm,nadd real*8 ttim, dt, c1, c2, c3, c4, c5, chi, dtcr integer idyn0 ttim,dt,c1,c2,c3,c4,c5, chi, dtcr, idyn0 common /tdata/ integer np ,up common /pointer/ np(400),up(200) real*8 hr integer mr / hr(1),mr(1) common /

Figure 5.2: FEAP Element Common Blocks.

include	'bdata.h'
include	'cdata.h'
include	'counts.h'
include	'eldata.h'
include	'elplot.h'
include	'eltran.h'
include	'hdata.h'
include	'iofile.h'
include	'prstrs.h'
include	'tdata.h'
include	'pointer.h'
include	'comblk.h'

Figure 5.3: FEAP Element Common Blocks using Includes.

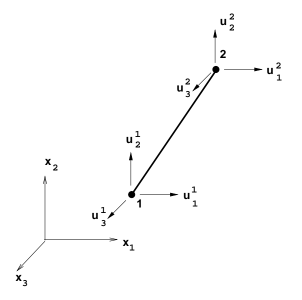


Figure 5.4: 2-Node Truss Element

```
if(isw.eq.3 .or. isw.eq.6) then
с
        Compute element length
        L2= 0.0d0
        do i = 1, ndm
         L2 = L2 + (xl(i,2) - xl(i,1))**2
        end do
       L = sqrt(L2)
        Compute strain-displacement matrix
С
        Lr = 1.d0/L2
        eps = 0.0d0
        do i = 1, ndm
         bb(i,1) = -(xl(i,2) - xl(i,1))*Lr
          bb(i,2) = -bb(i,1)
                = eps + bb(i,2)*(ul(i,2,1) - ul(i,1,1))
          eps
        end do
с
        Compute mass terms
        cmd = rhoA*L/3.0d0
        cmo = cmd * 0.5d0
       Form body/inertia force vector (dm = prop. ld.)
С
        sigA = EA*eps*L
        body = 0.5d0*L*dm
        do i = 1,ndm
         r(i,1) = body*d(6+i) - bb(i,1)*sigA
                - cmd*ul(i,1,5) - cmo*ul(i,2,5)
    &
          r(i,2) = body*d(6+i) - bb(i,2)*sigA
                 - cmo*ul(i,1,5) - cmd*ul(i,2,5)
     &
        end do
```

Figure 5.5: Element residual for two node truss

```
if(isw.eq.3) then
С
        Compute element length
        L2= 0.0d0
        do i = 1,ndm
          L2 = L2 + (xl(i,2) - xl(i,1))**2
        end do
        L = sqrt(L2)
        Form stiffness multiplier
С
        dd = ctan(1) * EA * L
        Compute strain-displacement matrix
С
        Lr = 1.d0/L2
        do i = 1, ndm
          bb(i,1) = -(xl(i,2) - xl(i,1))*Lr
          bb(i,2) = -bb(i,1)
db(i,1) = dd*bb(i,1)
          db(i,2) = -db(i,1)
        end do
```

Figure 5.6: Truss Tangent Matrix. Part 1

```
с
      Compute stiffness terms (N.B. ndm < or = ndf)
      i1 = 0
      do ii = 1,2
        j1 = 0
        do jj = 1,2
          do i = 1,ndm
            do j = 1,ndm
              s(i+i1, j+j1) = db(i, ii)*bb(j, jj)
            end do
          end do
        j1 = j1 + ndf
        end do
        i1 = i1 + ndf
      end do
      Compute mass terms and correct for inertial effects
с
      cmd = ctan(3)*rhoA*L/3.0d0
      cmo = cmd * 0.5d0
      do i = 1, ndm
        j = i + ndf
        s(i,i) = s(i,i) + cmd
        s(i,j) = s(i,j) + cmo
        s(j,i) = s(j,i) + cmo
        s(j,j) = s(j,j) + cmd
      end do
    endif
```

Figure 5.7: Truss Tangent Matrix. Part 2

```
subroutine slcn2d(sig,shp,xsj,sg,lint,nel,nes, p,s)
c-----[--.--+----.-------]
       Purpose: Project element variables to nodes
С
       Inputs:
С
С
         sig(nes,*) - Stresses at quadrature points
         shp(nel,*) - Shape functions at quadrature points
С
         xsj(*) - Volume element at quadrature points
sg(3,*) - Gauss points (1,2) and weights (3)
lint - Number of quadrature points
nel - Number nodes on element
С
С
С
       nel - Number nodes on element
nes - Dimension of stress array
С
С
  Outputs:
С
         p(nen) - Weights for 'lumped' projection
С
         s(nen,*) - Integral of variables
С
c-----[--.--+----.---+-----]
      implicit none
      include 'cdata.h' ! Contains 'nen'
      include 'strnum.h' ! Contains 'iste'
      integer i,l,lint,nel,nes
                xg,p(*),s(nen,*),xsj(*),sig(nes,*),shp(nel,*),sg(3,9)
      real*8
      do l = 1, lint
        do i = 1,nel
          xg = shp(i,l)*xsj(l)
p(i) = p(i) + xg
          s(i,1) = s(i,1) + sig(1,1)*xg
          s(i,2) = s(i,2) + sig(2,1)*xg
          s(i,3) = s(i,3) + sig(3,1)*xg
          s(i,4) = s(i,4) + sig(4,1) * xg
        end do ! i
      end do ! i
      iste = 4 ! Returns number projections
      end
```

Figure 5.8: Element variable projection routine

Chapter 6

UTILITY ROUTINES

The *FEAP* system includes several subprograms that can assist developers in writing new modules. In the next sections we describe some of the routines which perform numerical integration, compute shape functions and their derivatives, etc.

6.1 Numerical quadrature routines

Details on quadrature formula types and the layout and location of points and weights may be found in standard references.^[4, 5, 6, 2, 3] Here only the description of subroutine calls is included together with the available options on number of points.

6.1.1 One dimensional quadrature

Line integrals may be evaluated using Gaussian quadrature in which the approximation to an integral is given as

$$\int_{-1}^{+1} f(\xi) \,\mathrm{d}\xi \approx \sum_{l=1}^{L} f(\xi_l) \,W_l \tag{6.1}$$

where ξ_l are quadrature *points* and W_l are the *weights* to be applied at each point. The weights satisfy the condition.

$$\sum_{l=1}^{L} W_l = 2 . (6.2)$$

The Gauss-Legendre formula has points $|\xi_l|$ which are all less than unity. The subprogram call

CALL INT1D (L , SG , WG)

in which L is assigned an integer value between 1 and 5 returns the points in the array SG(*) and the weights in WG(*), both of which are of type REAL*8. The Gauss-Legendre formula integrates exactly polynomials up to order 2*L - 1.

The Gauss-Lobato formula has two of its points at -1 and 1 with the remainder in the interior of the interval. A routine to perform quadrature is obtain by using the call

```
CALL INT1DL ( L , SW )
```

in which L is assigned an integer value between 1 and 6. The values of the points and weights are returned in the two dimensional array SW: Points in SW(1,*) and weights in SW(2,*).

6.1.2 Two dimensional quadrature

Two dimensional quadrature on quadrilateral domains may be performed by repeated one-dimensional integration. The two dimensional integrations are approximated by

$$\iint_{-1}^{+1} f(\xi,\eta) \,\mathrm{d}\xi \,\mathrm{d}\eta \approx \sum_{l=1}^{L} f(\xi_l,\eta_l) \,W_l \tag{6.3}$$

where L is the total of all quadrature points. A routine to compute $n \times n$ order Gauss-Legendre quadrature is obtained by the call

```
CALL INT2D ( L , LINT, SW )
```

where L is assigned to the number of points in *each direction*, LINT is returned as the total number of points and SW(3,*) is an array containing the points and weights according to: SW(1,1) contains values of the points ξ_l ; SW(2,1) contains values of the points η_l ; and SW(3,1) contains values of the weights W_l .

Two dimensional quadrature on triangles may be performed using the subprograms call

Type	Number	Location
	Points	
1	1	Centroid $(O(h^2))$
3	3	Mid-sides $(O(h^3))$
-3	3	Interior $(O(h^3))$
4	4	Interior $(O(h^4))$ - Negative Wt.
6	6	Nodal $(O(h^3))$
-6	6	Interior $(O(h^4))$
7	7	Interior $(O(h^6))$
-7	7	Nodal $(O(h^4))$
12	12	Interior $(O(h^7))$
13	13	Interior $(O(h^8))$ - Negative Wt.

Table 6.1: Quadrature for triangles

CALL TINT2D (L , LINT, SW)

where L is a type indicator, LINT returns the number of points, and SW(4,*) is an array which returns three area coordinates and the quadrature weight: SW(1,1) returns the area coordinate L_{1l} (as defined in [2, 3]); SW(2,1) returns the area coordinate L_{2l} ; SW(3,1) returns the area coordinate L_{3l} ; SW(4,1) returns the weight W_l ; Table 6.1 describes the admissible types, number and location of quadrature points.

6.1.3 Three dimensional quadrature

Three dimensional quadrature on brick domains may be performed by repeated onedimensional integration. The three dimensional integrations are approximated by

$$\iiint_{-1}^{+1} f(\xi,\eta,\zeta) \,\mathrm{d}\xi \,\mathrm{d}\eta \,\mathrm{d}\zeta \approx \sum_{l=1}^{L} f(\xi_l,\eta_l,\zeta) \,W_l \tag{6.4}$$

where L is the total of all quadrature points. A routine to compute $n \times n \times n$ order Gauss-Legendre quadrature is obtained by the call

CALL INT3D (L , LINT, SW)

where L is assigned to the number of points in *each direction*, LINT is returned as the total number of points and SW(4,*) is an array containing the points and weights according to: SW(1,1) contains values of the points ξ_l ; SW(2,1) contains values of the

Type	Number	Location
	Points	
1	1	Centroid $(O(h^2))$
-1	4	Nodal $()(h^2))$
2	4	Interior $(O(h^3))$
3	5	Interior $(O(h^4))$ - Negative wt.
4	11	Interior $(O(h^4))$ - Negative wt.
-4	11	Nodal $(O(h^3))$
5	14	Interior $(O(h^5))$
6	16	Interior $(O(h^5))$
8	8	Node & Face $(O(h^2))$

Table 6.2: Quadrature for tetrahedra

points η_l ; and SW(3,1) contains values of the points ζ_l ; and SW(4,1) contains values of the weights W_l .

Three dimensional quadrature on tetrahedra may be performed using the subprograms call

CALL TINT3D (L , LINT, SW)

where L is a type indicator, LINT returns the number of points, and SW(5,*) is an array which returns three area coordinates and the quadrature weight: SW(1,1) returns the volume coordinate $L_{1,l}$ (as defined in [2,3]); SW(2,1) returns the volume coordinate $L_{2,l}$; SW(3,1) returns the volume coordinate $L_{3,l}$; SW(4,1) returns the volume coordinate $L_{4,l}$; SW(5,1) returns the weight W_l ; Table 6.2 describes the admissible types, number and location of quadrature points.

6.2 Shape function subprograms

Finite element approximations commonly use shape function subprograms to perform computations of the functions and their derivatives at preselected points (often the quadrature points). *FEAP* includes options to obtain the shape functions for some low order elements (linear and quadratic order) in one and two dimensions and linear shape functions for three dimensions. In addition a cubic Hermitian interpolation routine is available. The calling arguments for routines is summarized below.

6.2.1 Shape functions in one-dimension

Lagrangian interpolation in one-dimensional isoparametric forms may be obtained using the call

CALL SHP1D (S , XL , SHP, NDM, NEL, XJAC)

where

Parameter	Description
S	Natural coordinate ξ
XL(NDM,*)	Nodal coordinates for element
NDM	Spatial dimension of mesh
NEL	Number element nodes $(2 \text{ or } 3)$
SHP(2,NEL)	Shape function and derivative
XJAC	Jacobian transformation

The shape functions are evaluated as: SHP(1,i shape function derivative along the axis of the element and SHP(2,i) the shape function N_i . In calculations integrals are represented as

$$\int_{L} f(N_{i}, N_{i,s}) \,\mathrm{d}s = \int_{-1}^{1} f[N_{i}(\xi), N_{i,s}(\xi)] \, X JAC(\xi) \,\mathrm{d}\xi \tag{6.5}$$

and quadrature may be used for evaluation.

Calculation of natural coordinate derivatives only may be obtained with the call

CALL SHAP1DN(S , SHP, NEL)

where

Parameter	Description
S	Natural coordinate ξ
SHP(2,NEL)	Shape function and derivative
NEL	Number element nodes $(2 \text{ or } 3)$

where SHP(1,i contains $N_{i,\xi}$ and SHP(2,i) the shape function N_i .

Cubic Hermitian interpolation (e.g., for use in straight linear beam elements) given by

$$w = N_1^w \,\bar{w}_1 + N_2^w \,\bar{w}_2 + N_1^\theta \,\bar{\theta}_1 + N_2^\theta \,\bar{\theta}_2 \tag{6.6}$$

is obtained using the call

CALL SHP1DH (S , LEN , SHPW, SHPT)

where

Parameter	Description	
S	Natural coordinate ξ	
LEN	Length of the element (2-node)	
SHPW(4,2)	Shape functions for w_i	
SHPT(4,2)	Shape functions for θ_i	

The arrays are evaluated as follows:

- 1. SHPW(1,i), SHPT(1,i) are first derivatives (e.g. $N_{i,x}$);
- 2. SHPW(2,i), SHPT(2,i) are second derivatives (e.g. $N_{i,xx}$);
- 3. SHPW(3,i), SHPT(3,i) are third derivatives (e.g. $N_{i,xxx}$); and
- 4. SHPW(4,i), SHPT(4,i) are shape functions (e.g. N_i).

6.2.2 Shape functions in two-dimensions

Two-dimensional C_0 isoparametric interpolation on quadrilaterals of linear, quadratic and cubic order may be obtained using the subprogram call

CALL SHP2D (SS, XL, SHP, XJAC, NDM, NEL, IX, FLG)

where

Parameter	Description	
SS(2)	Natural coordinates ξ , η	
XL(NDM,NEL)	Element coordinates in local order	
NDM	Spatial dimension mesh $(2 \text{ or } 3)$	
NEL	Number nodes on element (4-9, 12, 16)	
IX(NEL)	Element global node numbers	
FLG	Return $\xi - \eta$ derivatives if true or	
	x - y derivatives if false	
SHP(3,NEL)	Shape functions and derivatives	
XJAC	Jacobian transformation from $x - y$ to $\xi - \eta$.	

The array SHP stores the values in the order: SHP(1,i) derivative with respect to ξ or x; SHP(2,i) derivative with respect to η or y; SHP(3,i) shape function.

Two-dimensional C_0 isoparametric interpolation on triangles of linear, quadratic and cubic order may be obtained using the subprogram call

CALL TRISHP (SS, XL, NDM, IORD, XJAC, SHP)

where

Parameter	Description
SS(3)	Area coordinates L_1, L_2, L_3
XL(NDM,*)	Element coordinates in local order
NDM	Spatial dimension mesh $(2 \text{ or } 3)$
IORD	Order of interpolation $(1 = 3 - node, 2 = 6 - node, 3 = 7 - node, 4 = 1)$
	6-node + 3 bubble, $10 = 10$ -node cubic)
XJAC	Jacobian transformation from $x - y$ to $\xi - \eta$
SHP(3,NEL	Shape functions and derivatives

The array SHP stores the values in the order: SHP(1,i) derivative with respect to ξ or x; SHP(2,i) derivative with respect to η or y; SHP(3,i) shape function. The parameter IORD defines the order of interpolation. If it is 1 simple 3-node triangles with linear interpolation is returned; if 2 quadratic interpolation; if 3 the interpolation is generated plus a cubic bubble in the seventh function. Giving the IORD parameter as a negative returns hierarchical form for mid side nodes.

6.2.3 Shape functions in three-dimensions

Three-dimensional C_0 isoparametric interpolation on bricks of linear order (i.e., 8-node elements) may be obtained using the subprogram call

CALL SHP3D (SS, XJAC, SHP, XL, NDM, NEL)

where

Parameter	Description		
SS(3)	Natural coordinates ξ , η , ζ		
XL(NDM,8)	Element coordinates in local order		
NDM	Spatial dimension mesh (2 or 3)		
NEL	Number nodes on element: $8 =$ linear brick; $20 =$ serendipity		
	quadratic; $27 = \text{Lagrangian quadratic}; 64 = \text{Lagrangian cubic}$		
SHP(4,8)	Shape functions and derivatives		
XJAC	Jacobian transformation from xyz to $\xi\eta\zeta$.		

The array SHP stores the values in the order: SHP(1,i) derivative with respect to x; SHP(2,i) derivative with respect to y; SHP(3,i) derivative with respect to z; SHP(4,i) shape function.

Three-dimensional C_0 isoparametric interpolation on tetrahedra of linear order (i.e., 4-node elements) may be obtained using the subprogram call

CALL TETSHP (SS, XL, NDM, NEL, XJAC, SHP)

where

Parameter	Description
SS(4)	Volume coordinates L_1, L_2, L_3, L_4
XL(NDM,4)	Element coordinates in local order
NDM	Spatial dimension mesh (3)
NEL	Number of nodes on element $(4, 10, 11, 14, 15)$
XJAC	Jacobian transformation from xyz to $\xi\eta\zeta$
SHP(4,4	Shape functions and derivatives

The array SHP stores the values in the same order as for the brick element.

6.3 Eigenvalues for 3×3 matrix

Three dimensional problems often require the solution of a 3×3 eigenproblem to generate principal values and directions. *FEAP* includes a special routine to calculate the values and vectors for symmetric arrays. The routine is used by a call to the subprogram as

CALL EIG3 (V, D, ROT)

On call to the routine V(3,3) is a REAL*8 array containing the symmetric array to be diagonalized. On return the eigenvalues are contained in D(3) and the vectors for each value in the columns of the V array. A Jacobi method is used with ROT an integer parameter returning the number of rotations to diagonalize. The routine is quite efficient compared to any attempt to compute vectors after closed form solution of the cubic for roots.

In addition to the general eigensolution above FEAP includes options to compute principal values of a symmetric second order tensor for two and three dimensional problems. In two dimensional use, the call to

CALL PSTR2D (SIG, PV)

is used where SIG(4) stores stresses in the order σ_{11} , σ_{22} , σ_{33} , σ_{12} and returns principal values and directions in PV(3) in the order σ_1 , σ_2 , and θ , where the angle is in degrees between x and the 1-axis. This routine does not use SIG(3).

In three dimensions the principal values are obtained using the call

CALL PSTR3D (SIG, PV)

where SIG(6) stores stresses in the order σ_{11} , σ_{22} , σ_{33} , σ_{12} , σ_{23} , σ_{31} , and returns principal values in PV(3) in the order σ_1 , σ_2 , σ_3 . Roots are ordered from most positive to most negative.

6.4 Plot routines

Several options exist in the FEAP system to create graphical plots for data and results.

6.4.1 Mesh plots

FEAP has plot capabilities to represent some standard element shapes. For continuum elements where the shape of the element is identical to the space dimension of the mesh (i.e., NDM) it is not necessary to provide any extra descriptions. However, if the dimension of the element topology is different from the mesh dimension it is necessary to add the include statement

include 'eldata.h'

and the statement

pstyp = pdim

within the ISW = 1 part of the element routine. Here pdim is the *dimension* of the element topology. For example, in a three dimensional shell problem where NDM = 3 and the element topology is two dimensional the statement is given as

$$pstyp = 2$$

Provided the nodal numbering of an element is as described in the *FEAP* User manual (i.e., numbered with vertex nodes first, followed by mid-side nodes, then face nodes and finally internal nodes) the program can use the actual number of nodes on the element to draw each element.

Failure to include a **pstyp** statement will usually result in unpredictable plots of the mesh and contour values.

The known types of plots are:

1. <u>Point element</u> with one node obtained by call

CALL PLTPT1 (IEL)

2. <u>Line element</u> with two nodes obtained by call

CALL PLTLN2 (IEL)

and for three node elements

CALL PLTLN3 (IEL)

3. Triangular element with 3-nodes obtained by call

CALL PLTRI3 (IEL)

and for 6-nodes obtained by call

CALL PLTRI6 (IEL)

4. Quadrilateral element with 4-nodes obtained by call

CALL PLQUD4 (IEL)

for 8- or 9-node elements the plot call is

CALL PLQUD8 (IEL)

and for 12- or 16-node quadrilaterals the call is

CALL PLTQ16 (IEL)

5. <u>Tetrahedral element</u> with 4-nodes obtained by call

CALL PLTET4 (IEL)

and for 10-node tetrahedra the call is

CALL PLTET10(IEL)

6. <u>Brick element</u> with 8-nodes obtained by call

CALL PLBRK8 (IEL)

and for 20- or 27-node bricks the call is

CALL PLBRK27(IEL)

Using these and internal extraction of element surfaces the program is able to make some hidden surface plots in three dimensions.

6.4.2 Element data plots

Users may construct plots within their elements (i.e., an ELMTnn) and access using the plot command:

PLOT, PELE, v1, v2, v3

In interactive mode in the plot environment it is only necessary to enter

PELE,v1,v2,v3

The values entered in v1, v2, v3 are optional and are passed to the element through a common block as

REAL*8 ELPLT COMMON /ELPDAT/ ELPLT(3)

The PELE option calls each element with the switch parameter ISW = 20. Users merely code whatever option they wish to include within their element module.

The standard color table is available through use of the subroutine call

CALL PPPCOL(ICOL, 0)

in which ICOL designates the color to be assigned according to Table 6.3. An exception occurs for PostScript outputs where black and white are switched (since the background then is assumed to be white).

ICOL	COLOR	ICOL	COLOR
0	Black	10	Green-yellow
1	White	11	Wheat
2	Red	12	Royal blue
3	Green	13	Purple
4	Blue	14	Aquamarine
5	Yellow	15	Violet-red
6	Cyan	16	Dark slate blue
7	Magenta	17	Grey
8	Orange	18	Light grey
9	Coral		

Table 6.3: Color pallet for FEAP plots

A straight line segment may be drawn to the screen in the current color between the coordinates (x_1, y_1, z_1) and (x_2, y_2, z_2) using the commands

CALL PLOTL(X1,Y1,Z1, 3) CALL PLOTL(X1,Y1,Z1, 2)

Here the basic command is

IP	Action
1	Start panel fill
2	Move to point
3	Draw to point

Table 6.4: Values for control of plots

CALL PLOTL(X1,Y1,Z1, IP)

where the three cartesian coordinates relate to mesh coordinates (not screen values) and IP is a parameter defined according to Table 6.4.

The perimeter of a panel is drawn with standard line drawing commands starting with

CALL PLOTL(X1,Y1,Z1, 1)

and continuing with a sequence of draw commands

CALL PLOTL(Xi,Yi,Zi, 2)

(however, no lines appear on the screen) and the fill of each panel is completed by the statement

CALL CLPAN

It should be noted that all plots within FEAP are performed in three dimensions. For two dimensional problems no z_i coordinates are available in the XL(NDM,NEN) array and, hence, it is necessary to assign zero values for the z_i coordinates before calling a plot subprogram. If a perspective view has been requested a full use of a x_i, y_i, z_i specification is made. In this case a user may wish to pass the value of some solution variable as the z_i value (scaled so that it will make sense relative to the x_i, y_i coordinate values). Similarly, if deformed plots are being performed it is necessary to add (scaled) displacements to the coordinates. The current value of the scaling parameter (i.e., variable CS) is available in labeled common PVIEW. In this case one can add the statements (assuming here that the displacements correspond to the coordinate directions)

DO NE = 1, NEL

```
DO I = 1,NDM
XP(I,NE) = XL(I,NE) + CS*UL(I,NE)
END DO ! I
END DO ! NE
```

(NEL is the number of connected nodes to each element and is passed through labeled common ELDATA) before performing any deformed plots and then plot the appropriate values of XP. Indeed, this may always be performed as the value of CS will be zero for an *undeformed* plot.

6.4.3 Other user plots

It is also possible for users to prepare plot outputs unrelated to elements. The plot command

PLOT UPLOt v1 v2 v3

initiates a call to the subroutine UPLOT which has the basic structure

```
SUBROUTINE UPLOT(CT)
IMPLICIT NONE
REAL*8 CT(3)
...
END
```

The argument CT contains the values for the three parameters v1, v2, v3. The default color is *white*. Direct plots in screen coordinates [lower left at (0,0); upper right at (1,1)] may be given using the statement

CALL DPLOT(XS, YS, IP)

where XS, YS are between zero (0) and one (1) and IP is interpreted according to Table 6.4. Panels are closed using

```
CALL CLPAN
```

and colors treated according to values specified in calls to PPPCOL.

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