

*User's Manual for the FEHM  
Application — A Finite-Element  
Heat- and Mass-Transfer Code*

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*User's Manual for the FEHM  
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# **User's Manual for the FEHM Application— A Finite-Element Heat- and Mass-Transfer Code**

by

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## **ABSTRACT**

This document is a manual for the use of the FEHM application, a finite-element heat- and mass-transfer computer code that can simulate nonisothermal multiphase multicomponent flow in porous media. The use of this code is applicable to natural-state studies of geothermal systems and groundwater flow. A primary use of the FEHM application will be to assist in the understanding of flow fields and mass transport in the saturated and unsaturated zones below the proposed Yucca Mountain nuclear waste repository in Nevada. The equations of heat and mass transfer for multiphase flow in porous and permeable media are solved in the FEHM application by using the finite-element method. The permeability and porosity of the medium are allowed to depend on pressure and temperature. The code also has provisions for movable air and water phases and noncoupled tracers; that is, tracer solutions that do not affect the heat- and mass-transfer solutions. The tracers can be passive or reactive. The code can simulate two-dimensional, two-dimensional radial, or three-dimensional geometries. In fact, FEHM is capable of describing flow that is dominated in many areas by fracture and fault flow, including the inherently three-dimensional flow that results from permeation to and from faults and fractures. The code can handle coupled heat and mass-transfer effects, such as boiling, dryout, and condensation that can occur in the near-field region surrounding the potential repository and the natural convection that occurs through Yucca Mountain due to seasonal temperature changes. The code is also capable of incorporating the various adsorption mechanisms, ranging from simple linear relations to nonlinear isotherms, needed to describe the very complex transport processes at Yucca Mountain. This report outlines the uses and capabilities of the FEHM application, initialization of code variables, restart procedures, and error processing. The report describes all the data files, the input data, including individual input records or parameters, and the various output files. The system interface is described, including the software environment and installation instructions. Examples illustrating various aspects of the code are sprinkled throughout the report, and the final section demonstrates how to construct an input file, shows typical code execution, and gives three examples: heat conduction in a square, a reactive-transport problem, and Problem 5, Case A, of the DOE Code Comparison Project.

## 1.0 PURPOSE

This User's Manual documents the use of the finite-element heat- and mass-transfer (FEHM) application (Zyvoloski, et al. 1988).

## 2.0 DEFINITIONS AND ACRONYMS

### 2.1 Definitions

FEHM: finite-element heat- and mass-transfer code.

FEHMN: an earlier version of FEHM designed specifically for the Yucca Mountain Site Characterization Project. Both versions are now equivalent, and the use of FEHMN has been dropped.

### 2.2 Acronyms

AVS: Advanced Visual Systems.

I/O: input/output.

LANL: Los Alamos National Laboratory.

UCD: unstructured cell data.

YMP: Yucca Mountain Site Characterization Project.

## 3.0 REFERENCES

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## 4.0 PROGRAM CONSIDERATIONS

### 4.1 Program Options

The uses and capabilities of FEHM are summarized in Table I with reference to the macro input structure discussed in Section 6.0.

**Table I. Capabilities of FEHM with macro command references**

- |      |   |
|------|---|
| I.   | Mass and energy balances in porous media  |
| A.   | Variable rock properties ( <b>rock</b> )  |
| B.   | Variable permeability ( <b>perm</b> )   |
| C.   | Variable thermal conductivity ( <b>cond</b> )   |
| D.   | Variable fracture properties, dual porosity, and dual porosity/dual permeability ( <b>dual, dpdp</b> )              |
| II.  | Multiple components available   |
| A.   | Air-water isothermal mixture available ( <b>airwater</b> ), fully coupled to heat and mass transfer ( <b>ngas</b> ) |
| B.   | Up to 10 solutes with chemical reactions between each ( <b>trac, rxn</b> )  |
| C.   | Several different capillary pressure models ( <b>cap</b> )  |
| D.   | Several different relative permeability models ( <b>rlp</b> )   |
| III. | Equation of state flexibility inherent in code ( <b>eos</b> )   |
| IV.  | Pseudo-stress models available  |
| A.   | Linear porosity deformation ( <b>ppor</b> )   |
| B.   | Gangi stress model ( <b>ppor</b> )  |
| V.   | Numerics  |
| A.   | Finite element with multiple element capabilities ( <b>elem</b> )   |
| B.   | Short form input methods available ( <b>coor, elem</b> )  |
| C.   | Flexible properties assignment ( <b>zone</b> )  |
| D.   | Flexible solution methods   |
| 1.   | Upwinding, implicit solution available ( <b>ctrl</b> )  |
| 2.   | Iteration control adaptive strategy ( <b>iter</b> )   |
| E.   | Finite volume geometry ( <b>finv</b> )  |
| VI.  | Flexible time step and stability control ( <b>time</b> )  |

### 4.2 Initialization

The coefficient arrays for the polynomial representations of the density (**crl, crv**), enthalpy (**cel, cev**), and viscosity (**cvl, cvv**) functions are initialized to the values enumerated in the FEHM document "Models and Methods Summary" (Zyvoloski et al. 1997, Table III). Values for the saturation pressure and temperature function coefficients are also found in that document (Table IV). All other global array and scalar variables, with the exception of the variables listed below in Table II, whether integer or real, are initialized to zero.

Table II. Initial (default) values					
Variable	Value	Variable	Value	Variable	Value
aiaa	1.0	contim	1.0e+30	daymin	1.0e-05
daymax	30.0	g1	1.0e-06	g2	1.0e-06
g3	1.0e-03	iamx	500	ncntr	10000000
nicg	1	rnmax	1.0e+11	str	1.0
strd	1.0	tmch	1.0e-09	tmelt	-1.0e+12
upwgt	1.0	upwgt <sub>a</sub>	1.0		

### 4.3 Restart Procedures

FEHM writes a restart file for each run. The name of the restart output file may be given in the input control file or as terminal input, or if unspecified, it will default to *fehmn.fin* (see Section 6.2.1). The file is used on a subsequent run by providing the name of the generated file (via control file or terminal) for the name of the restart input file. It is recommended that the name of the restart input file be modified to avoid confusion with the restart output file. For example, by changing the suffix to *.ini*, the default restart output file, *fehmn.fin*, would be renamed *fehmn.ini* and that file name placed in the control file or given as terminal input. Values from the restart file will overwrite any variable initialization prescribed in the input file. The initial time of simulation will also be taken from the restart file.

### 4.4 Error Processing

Due to the nonlinearity of the underlying partial differential equations, it is possible to produce an underflow or overflow condition through an unphysical choice of input parameters. More likely, the code will fail to converge or will produce results that are out of bounds for the thermodynamic functions. The code will attempt to decrease the time step until convergence occurs. If the time step drops below a prescribed minimum, the code will stop, writing a restart file. The user is encouraged to look at the input check file, which contains information regarding maximum and minimum values of key variables in the code. All error and warning messages will be sent to an output error file.

Table III provides additional information on errors that will cause FEHM to terminate.

**Table III. Error conditions that result in program termination**

Error condition	Error message
<b>I/O file error</b>	
Unable to open I/O file	<pre> **** Error opening file <i>fileid</i> **** . . . ****-----**** ****          JOB STOPPED          **** ****-----**** </pre>
Coefficient storage file not found	program terminated because coefficient storage file not found
Optional rlp file not found	relative perm file does not exist: stopping
Optional input file not found	ERROR nonexistent file <i>filename</i> STOPPED trying to use optional input file
Unable to open optional input file	ERROR opening <i>filename</i> STOPPED trying to use optional input file
Unable to determine file prefix for AVS output files	FILE ERROR: nmfil2 file: <i>filename</i> unable to determine contour file prefix
<b>Input deck errors</b>	
Coordinate or element data not found	<pre> ****      COOR Required Input      **** -or- ****      ELEM Required Input      ****  ****-----**** ****          JOB STOPPED          **** ****-----**** </pre>
Invalid macro read	**** error in input deck : <i>char</i> ****
Invalid AVS keyword read for macro <b>cont</b>	<pre> ERROR:READ_AVS_IO unexpected character string (terminate program execution) Valid options are shown: . . . The invalid string was: <i>string</i> </pre>

Table III. Error conditions that result in program termination (continued)	
Error condition	Error message
Invalid parameter values (macros using loop construct)	Fatal error - for array number <i>arraynum</i> macro - <i>macro</i> Group number - <i>groupnum</i> Something other than a real or integer has been specified -or- Line number - <i>line</i> Bad input, check this line
Invalid tracer input	** Using Old Input Enter Temperature Dependency Model Number: 1 - Van Hoff 2 - awwa model, see manual for details **
Invalid transport conditions	Fatal error. You specified a Henrys Law species with initial concentrations input for the vapor phase ( <i>icns</i> = -2), yet the Henrys Constant is computed as 0 for species number <i>speciesnum</i> and node number <i>nodenum</i> . If you want to simulate a vapor- borne species with no interphase transport, then you must specify a gaseous species ( <i>icns</i> = -1).
Invalid flag specified for diffusion coefficient calculation	ERROR -- Illegal Flag to concadiff Code Aborted in concadiff
Optional input file contains data for wrong macro	ERROR --> Macro name in file for macro <i>macroname</i> is <i>wrong_macroname</i> STOPPED trying to use optional input file
<b>Invalid parameters set</b>	
Dual porosity	**** check fracture volumes, stopping**** **** check equivalent continuum VGs ****
Noncondensable gas	cannot input <i>ngas</i> temp in single phase -or- <i>ngas</i> pressure lt 0 at temp and total press given max allowable temperature <i>temp</i> -or- <i>ngas</i> pressure gt total pressure <i>i</i> = <i>i</i> -or- <i>ngas</i> pressure lt 0.
Particle tracking	ERROR: <i>Pcnsk</i> in <i>ptrk</i> must be either always positive or always negative. Code aborted in <i>set_ptrk.f</i>

Table III. Error conditions that result in program termination (continued)	
Error condition	Error message
Tracer	ERROR: Can not have both particle tracking (ptrk) and tracer input (trac). Code Aborted in concn.f
<b>Insufficient storage</b>	
Geometric coefficients	program terminated because of insufficient storage
Dual porosity	***** n > n0, stopping *****
<b>Too many negative volumes or finite-element coefficients</b>	too many negative volumes:stopping -or- too many negative coefficients:stopping
<b>Unable to compute local coordinates</b>	iteration in zone did not converge, izeone = zone please check icnl in macro CTRL
<b>Singular matrix in LU decomposition</b>	singular matrix in ludcmp
<b>Solution failed to converge</b>	timestep less than daymin timestep_number current_timestep_size current_simulation_time -or- Tracer Time Step Smaller Than Minimum Step Stop in resetttrc



## 5.0 DATA FILES

### 5.1 Control File (iocntl)

#### 5.1.1 Content

The control file contains the names of the input and output files needed by the FEHM code. In addition to listing the I/O file names, the terminal (tty) output option and the user subroutine number are given. The control file provides the user an alternate means for inputting file names, terminal output option, and user subroutine number than through the terminal I/O. It is useful when long file names are used or when files are buried in several subdirectories, and it is also useful for automated program execution.

#### 5.1.2 Use by program

The control file is an input file that provides the FEHM application with the names of the input and output files, the terminal output units, and the user subroutine number to be used for a particular run. The default control file name is *fehmn.files*. If the control file is found, it is read prior to problem initialization. If not present, terminal I/O is initiated, and the user is prompted for required information. A control file may use a name other than the default. This alternate control file name would be input during terminal I/O (see Section 6.1.1.1).

#### 5.1.3 Auxiliary processing

N/A

### 5.2 Input File (inpt)

#### 5.2.1 Content

The input file contains user-parameter initialization values and problem-control information. The form of the file name is *filen* or *filen.\**, where "*filen*" is a prefix used by the code to name auxiliary files and "\*" represents an arbitrary file extension. If a file name is not specified when requested during terminal I/O, the file *fehmn.dat* is the default. The organization of the file is described in detail in Section 6.2.

#### 5.2.2 Use by program

The input file is an input file that provides the FEHM application with user-parameter initialization values and problem-control information. The input file is read during problem initialization.

#### 5.2.3 Auxiliary processing

N/A

## **5.3 Geometry Data File (incoor)**

### **5.3.1 Content**

The geometry data file contains the mesh-element and coordinate data. This file can either be the same as the input file or a separate file.

### **5.3.2 Use by program**

The geometry data file is an input file that provides the FEHM application with element and coordinate data. The geometry data file is read during problem initialization.

### **5.3.3 Auxiliary processing**

N/A

## **5.4 Zone Data File (inzone)**

### **5.4.1 Content**

The zone data file contains the zone information (see macro **zone**). This file can either be the same as the input file or a separate file.

### **5.4.2 Use by program**

The zone data file is an input file that provides the FEHM application with geometric-zone descriptions. The zone data file is read during problem initialization.

### **5.4.3 Auxiliary processing**

N/A

## **5.5 Optional Input Files**

### **5.5.1 Content**

The optional input files contain user-parameter initialization values and problem-control information. The names of optional input files are provided in the main input file to direct the code to auxiliary files to be used for data input. Their use is described in detail in Section 6.2.

### **5.5.2 Use by program**

The optional input file is an auxiliary input file that provides the FEHM application with user-parameter initialization values and problem-control information. The optional input files are read during problem initialization.

### **5.5.3 Auxiliary processing**

N/A

## **5.6 Output File (iout)**

### **5.6.1 Content**

The output file contains the FEHM output. The file name is provided in the input control file or as terminal input, or it may be generated by the code from the name of the input file if terminal I/O is invoked. The generated name is of the form *filen.out*, where the "filen" prefix is common to the input file.

### **5.6.2 Use by program**

The output file is an output file the FEHM application uses for general program time-step summary information. It is accessed throughout the program as the simulation steps through time.

### **5.6.3 Auxiliary processing**

N/A

## **5.7 Read File (iread)**

### **5.7.1 Content**

The read file contains the initial values of pressure, temperature, saturation, and simulation time (the restart or initial state values). The naming convention is similar to that for the output file. The generated name is of the form *filen.ini*.

### **5.7.2 Use by program**

The read file is an input file the FEHM application uses for program restarts. The read file is read during problem initialization.

### **5.7.3 Auxiliary processing**

N/A

## **5.8 Write File (isave)**

### **5.8.1 Content**

The write file contains the final values of pressure, temperature, saturation, and simulation time for the run. This file can in turn be used as the read file in a restart run. The naming convention is similar to that for the output file. The generated name is of the form *filen.fin*.

### **5.8.2 Use by program**

The write file is an output file the FEHM application uses for storing state data of the simulation. It is accessed at specified times throughout the program when state data should be stored.

### **5.8.3 Auxiliary processing**

N/A

## 5.9 History Plot File (ishis)

### 5.9.1 Content

The history plot file contains data for time history plots of variables. The naming convention is similar to that for the output file. The generated name is of the form *filen.his*.

### 5.9.2 Use by program

The history plot file is an output file the FEHM application uses for storing time history data for pressure, temperature, flow, and energy output. It is accessed throughout the program as the simulation steps through time.

### 5.9.3 Auxiliary processing

This file is used to produce time history plots with the Browser (see Section 8.5).

## 5.10 Solute Plot File (istrc)

### 5.10.1 Content

The solute plot file contains time history data for solute concentrations at specified nodes. The naming convention is similar to that for the output file. The generated name is of the form *filen.trc*.

### 5.10.2 Use by program

The solute plot file is an output file the FEHM application uses for storing time history data for tracer output. It is accessed throughout the program as the simulation steps through time.

### 5.10.3 Auxiliary processing

This file is used to produce time history plots of tracers with the Browser (see Section 8.5).

## 5.11 Contour Plot File (iscon)

### 5.11.1 Content

The contour plot file contains the contour plot data. The naming convention is similar to that for the output file. The generated name is of the form *filen.con*.

### 5.11.2 Use by program

The contour plot file is an output file the FEHM application uses for storing contour data for pressure, temperature, flow, energy output, and tracer output. It is accessed at specified times throughout the program when contour data should be stored.

### 5.11.3 Auxiliary processing

N/A

## 5.12 Contour Plot File for dual or dpdp (iscon1)

### 5.12.1 Content

The dual or dpdp contour plot file contains the contour plot data for dual-porosity or dual-porosity/dual-permeability problems. The naming convention is similar to that for the output file. The generated name is of the form *filen.dp*.

### 5.12.2 Use by program

The dual or dpdp contour plot file is an output file the FEHM application uses for storing contour data for pressure, temperature, flow, energy output, and tracer output for dual-porosity or dual-porosity/dual-permeability problems. It is accessed at specified times throughout the program when contour data should be stored.

### 5.12.3 Auxiliary processing

N/A

## 5.13 Stiffness Matrix Data File (isstor)

### 5.13.1 Content

The stiffness matrix data file contains finite-element coefficients calculated by the code. It is useful for repeated calculations that use the same mesh, especially for large problems. The naming convention is similar to that for the output file. The generated name is of the form *filen.stor*.

### 5.13.2 Use by program

The stiffness matrix data file is both an input and an output file that the FEHM application uses for storing or reading finite-element coefficients calculated by the code. The stiffness matrix data file is read during problem initialization, if being used for input. It is accessed after finite-element coefficients are calculated, if being used for output.

### 5.13.3 Auxiliary processing

N/A

## 5.14 Input Check File (ischk)

### 5.14.1 Content

The input check file contains a summary of coordinate and variable information, suggestions for reducing storage, coordinates at which maximum and minimum values occur, and information about input for variables set at each node. The naming convention is similar to that for the output file. The generated name is of the form *filen.chk*.

#### 5.14.2 Use by program

The input check file is an output file the FEHM application uses for writing a summary of the data initialization. The input check file is accessed after data initialization has been completed.

#### 5.14.3 Auxiliary processing

N/A

### 5.15 Output Error File (ierr)

#### 5.15.1 Content

The output error file contains any error or warning messages issued by the code during a run. The file is always named *fehmn.err* and will be found in the directory from which the problem was executed.

#### 5.15.2 Use by program

The output error file is an output file the FEHM application uses for writing error or warning messages issued by the code during a run. It may be accessed at any time.

#### 5.15.3 Auxiliary processing

N/A

### 5.16 Advanced Visual Systems (AVS) Output Files

#### 5.16.1 Content

The Advanced Visual Systems (AVS) output files contain geometry-based data that can be imported into AVS UCD (unstructured cell data) graphics routines. The AVS output files each have a unique file name indicating the section type, the data type, and the time step at which the files were created. These file names are automatically generated by the code and are of the form *fileprefix.NumberAVS\_id*, where *fileprefix* is common to the contour-output-file prefix if defined, otherwise, it is the input-file prefix; *Number* is a value between 10001 and 99999; and *AVS\_id* is a string denoting file content (see Table IV). In general, *\_head* are header files, *\_geo* is the geometry file, and *\_node* with *\_mat*, *\_sca*, *\_vec*, *\_con*, *\_mat\_dual*, *\_sca\_dual*, *\_vec\_dual*, or *\_con\_dual* are the data selected for output. Currently all properties are node-based rather than cell-based.

**Table IV. AVS file content tag**

<i>AVS_id</i>	File purpose
<i>_avs_log</i>	Log file from AVS output routines
<i>_geo</i>	Geometry output file containing coordinates and cell information
<i>_mat_head</i>	AVS UCD header for material properties file

<b>Table IV. AVS file content tag (continued)</b>	
<b>AVS_id</b>	<b>File purpose</b>
_mat_dual_head	AVS UCD header for material properties file for dual or dpdp
_sca_head	AVS UCD header for scalar-parameter values file
_sca_dual_head	AVS UCD header for scalar-parameter values file for dual or dpdp
_vec_head	AVS UCD header for vector-parameter values
_vec_dual_head	AVS UCD header for vector-parameter values for dual or dpdp
_con_head	AVS UCD header for solute concentration file
_con_dual_head	AVS UCD header for solute concentration file for dual or dpdp
_mat_node	Data output file with material properties
_mat_dual_node	Data output file with material properties for dual or dpdp
_sca_node	Data output file with scalar-parameter values (pressure, temperature, saturation)
_sca_dual_node	Data output file with scalar-parameter values (pressure, temperature, saturation) for dual or dpdp
_vec_node	Data output file with vector-parameter values (velocity)
_vec_dual_node	Data output file with vector-parameter values (velocity) for dual or dpdp
_con_node	Data output file with solute concentration
_con_dual_node	Data output file with solute concentration for dual or dpdp

### 5.16.2 Use by program

The AVS output files are output files the FEHM application uses for storing geometry-based data for material properties, temperature, saturation, pressure, velocities, and solute concentrations in a format readable by AVS graphics. The log output file is created on the first call to the AVS write routines. It includes the code version number, date, and problem title. When output for a specified time step has been completed, a line containing the file-name prefix, time step, call number (this variable is 1 for the initial call and is incremented with each call to write AVS contour data), and time (days) is written. The header files, one for each type of data being stored, and the single geometry file are written during the first call to the AVS output routines. The node data files are written for each

call to the AVS write routines at specified times throughout the program when contour data should be stored using AVS format.

### 5.16.3 Auxiliary processing

These files are used for visualization and analysis of data by AVS and to produce contour plots by the Browser (see Section 8.5).

To use these with AVS, the appropriate header file, geometry file, and data file for each node must be concatenated into one file of the form *filen.inp* (Fig. 1). This concatenation can be done with the script **fehm2avs** for a series of files with the same root *filen* or manually, for example:

```
cat filen.10001_head filen.10001_geo filen.10001_mat_node > filen.10001.inp
```

Once header and geometry have been merged with data files into a single AVS file, the data can be imported into AVS using the read\_uct module.

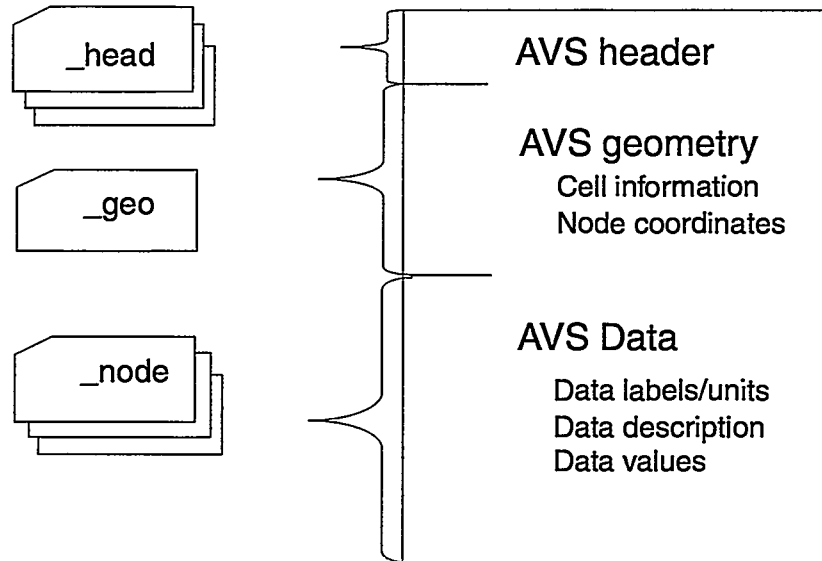


Figure 1. AVS UCD-formatted FEHM output files.



## 6.0 INPUT DATA

### 6.1 General Considerations

#### 6.1.1 Techniques

##### 6.1.1.1 Control file or terminal I/O startup

The input/output (I/O) file information is provided to the code from an input control file or the terminal. The default control file name is *fehmn.files*. If a control file with the default name is present in the directory from which the code is being executed, no terminal input is required. If the default control file is not present, input prompts are written to the screen preceded by a short description of the I/O files used by FEHM. The descriptions of the I/O files are elaborated on in Section 5.0. The initial prompt asks for the name of a control file. If a control file name is entered for that prompt, no further terminal input is required. If a control file is not used, the user is then prompted for I/O file names, the tty output flag, and user subroutine number. When the input file name is entered from the terminal, the user has the option of letting the code generate the names for the remainder of the auxiliary files using the input file-name prefix. The form of the input file name is *filen* or *filen.\**, where "*filen*" is the prefix used by the code to name the auxiliary files and ".\*" represents an arbitrary file extension.

##### 6.1.1.2 Macro control structure

The finite-element heat- and mass-transfer code (FEHM) contains a macro control structure for data input that offers added flexibility to the input process. The macro command structure makes use of a set of control statements recognized by the input module of the program. When a macro control statement is encountered in an input file, a certain set of data with a prescribed format is expected and read from the input file. In this way, the input is divided into separate, unordered blocks of data. The input file is therefore a collection of macro control statements, each followed by its associated data block. Blocks of data can be entered in any order, and any blocks unnecessary to a particular problem need not be entered. The macro control statements must appear in the first four columns of a line. The other entries are free format, which adds flexibility but requires that values be entered for all input variables (no assumed null values).

As an aid to the user, the capabilities of FEHM summarized in Table I refer to applicable macro commands. Table V lists the macro control statements with a brief description of the data associated with each. A more detailed description of each macro control statement and its associated input are found in

**Table V. Macro control statements for FEHM**

<b>Control statement</b>	<b>Description</b>
<b>adif</b>	Air-water vapor diffusion
<b>airwater</b>	Isothermal air-water input
<b>alti</b>	Alternate input
<b>boun</b>	Implement boundary conditions
<b>bous</b>	Boussinesq-type approximation
<b>cap</b>	No longer used, see macro rlp
<b>cond</b>	Thermal-conductivity data ( <b>required for nonisothermal problem</b> )
<b>cont</b>	Contour plot data
<b>coor</b>	Node-coordinate data ( <b>required</b> )
<b>ctrl</b>	Program-control parameters ( <b>required</b> )
<b>dof</b>	[Not implemented]
<b>dmdp</b>	Double-porosity/double-permeability model input
<b>dual</b>	Input for dual-porosity solution
<b>elem</b>	Element-node data ( <b>required</b> )
<b>eos</b>	Equation-of-state data
<b>exrl</b>	Explicit relative permeability
<b>finv</b>	Finite-volume flow coefficients
<b>flow</b>	Flow data
<b>flo2</b>	Alternate format for flow data
<b>flxo</b>	Flux printout
<b>head</b>	Hydraulic-head values
<b>hflx</b>	Heat-flux input
<b>ice</b>	Ice-phase calculations (untested)
<b>init</b>	Initial value data ( <b>required if macro pres or restart file is not used</b> )
<b>iter</b>	Iteration parameters
<b>itup</b>	Iterations used with upwinding
<b>iupk</b>	Upwind transmissibility including intrinsic permeability
<b>ivfc</b>	Enables volume-factor calculations
<b>mdnode</b>	Enables extra connections to be made to nodes

**Table V. Macro control statements for FEHM (continued)**

<b>Control statement</b>	<b>Description</b>
<b>ngas</b>	Noncondensable-gas (air) data
<b>nod2</b>	Node numbers for output and time histories, and alternate nodes for terminal output
<b>node</b>	Node numbers for output and time histories
<b>perm</b>	Permeability input ( <b>required</b> )
<b>pest</b>	Estimation routine
<b>ppor</b>	Pressure- and temperature-dependent porosity and permeability
<b>pres</b>	Initial pressure, temperature, and saturation data boundary-conditions specification ( <b>required if macro init or restart file is not used</b> )
<b>ptrk</b>	Particle-tracking simulation input
<b>renm</b>	Renumbers nodes
<b>rflx</b>	Radiation-flux input
<b>rlp</b>	Relative-permeability input ( <b>required for 2-phase problem, otherwise optional</b> )
<b>rock</b>	Rock-density, specific-heat, and porosity input ( <b>required</b> )
<b>rxn</b>	Chemical reaction-rate model input
<b>sol</b>	Solver specifications ( <b>required</b> )
<b>solv</b>	[Not implemented]
<b>stea</b>	Steady-state solution generated for initial variable field
<b>stop</b>	Signals the end of input ( <b>required</b> )
<b>strs</b>	Initiates stress solution [Not implemented in this version of FEHM]
<b>text</b>	Text input to be written to output file
<b>thic</b>	Variable-thickness input for two-dimensional problems
<b>time</b>	Time-step and time-of-simulation data ( <b>required</b> )
<b>trac</b>	Solute-simulation input
<b>user</b>	User subroutine call
<b>vcon</b>	Variable thermal-conductivity input
<b>velo</b>	Velocity printout
<b>wlbr</b>	Wellbore-style input
<b>zone</b>	Geometric definition of grid for input-parameter assignment

Section 6.2. Macro control statements may be called more than once, if for example, the user wishes to reset some property values after defining alternate zones. Some statements are required, as indicated in Table V; the others are optional.

Comments may be entered in the input file by beginning a line with a “#” symbol (the “#” symbol must be found in the first column of the line). Comments may precede or follow macro blocks but may not be found within a block.

Optional input files may be used by substituting a keyword and file name in the main input file (described in detail in Section 6.2.2). The normal macro input is then entered in the auxiliary file.

Many input parameters, such as porosity or permeability, vary throughout the grid and need to have different values assigned at different nodes. This task is accomplished in two ways. The first uses a nodal loop-type definition (which is the default):

**JA, JB, JC, PROP1, PROP2 ... ,**

where

**JA** - first node to be assigned with the properties PROP1, PROP2 ... ,

**JB** - last node to be assigned with the properties PROP1, PROP2 ... ,

**JC** - loop increment for assigning properties PROP1, PROP2 ... , and

PROP1, PROP2 - property values to be assigned to the indicated nodes.

In the input blocks using this structure, one or more properties are manually entered in the above structure. When a blank line is entered, that input block is terminated and the code proceeds to the next group or control statement. (Note that blank input lines are shaded in the examples shown in Section 6.2). The nodal definition above is useful in simple geometries for which the node numbers are easily found. Boundary nodes often come at regular node intervals, and the increment counter **JC** can be adjusted so the boundary conditions are easily entered. To set the same property values at every node, the user may set **JA** and **JC** to 1 and **JB** to the total number of nodes or, alternatively, set **JA** = 1 and **JB** = **JC** = 0.

For dual-porosity problems, which have three sets of parameter values at any nodal position, nodes 1 to N (where N is the total number of nodes in the grid (see macro **coor**)) represent the fracture nodes, nodes N + 1 to 2N are generated for the second set of nodes, the first matrix material, and nodes 2N + 1 to 3N for the third set of nodes, the second matrix material. For double-

porosity/double-permeability problems, which have two sets of parameter values at any nodal position, nodes 1 to N represent the fracture nodes and nodes N + 1 to 2N are generated for the matrix material.

For more complicated geometries, such as 3-D grids, the node numbers are often difficult to determine. Here a geometric description is preferred. To enable the geometric description, the **zone** control statement (page 74) is used in the input file before the other property macro statements occur. The input macro **zone** requires the specification of the coordinates of 4-node parallelograms for 2-D problems or 8-node polyhedrons in 3-D. In one usage of the control statement **zone**, all the nodes are placed in geometric zones and assigned an identifying number. This number is then addressed in the property input macro commands by specifying a **JA** < 0 in the definition of the loop parameters given above. For example, if **JA** = -1, the properties defined on the input line would be assigned to the nodes defined as belonging to geometric Zone 1 (**JB** and **JC** must be input but are ignored in this case). The control statement **zone** may be called multiple times to redefine geometric groupings for subsequent input. The previous zone definitions are not retained between calls. Up to 100 zones may be defined. For dual-porosity problems, which have three sets of parameter values at any nodal position, Zone 100 + I is the zone number for the second set of nodes defined by Zone I, and Zone 200 + I is the zone number for the third set of nodes defined by Zone I. For double-porosity/double-permeability problems, which have two sets of parameter values at any nodal position, Zone 100 + I is the zone number for the second set of nodes defined by Zone I.

### 6.1.2 Consecutive cases

The program retains no input data between cases. The values of all variables are reinitialized with each run, either from the input files or a restart file when used.

### 6.1.3 Defaults

Default values are set during the initialization process if overriding input is not provided by the user.

## 6.2 Individual Input Records or Parameters

Other than the control file or terminal I/O, the main user input is provided using macro control statements in the input file, geometry data file, or zone data file. Data provided in the input files are entered in free format with the exception of the macro control statements, which must appear in the first four columns of a line. Data values may be separated with spaces, commas, or tabs. The primary input file differs from the others in that it begins with a title line (80 characters

maximum) followed by input in the form of the macro commands. Each file containing macro commands should be terminated with the **stop** control statement. In the examples provided in the following subsections, blank input lines are depicted with shading.

### 6.2.1 Control file or terminal I/O input

The parameters enumerated below are entered in order, one per line, in the control file (excluding the control file name [nmfile(1)]) or as prompted for during terminal input. If there is a control file with the name *fehmn.files* in your local space, FEHM will execute using that control file, and there will be no prompts. If another name is used for the control file, it can be entered at the first prompt.

A blank line can be entered in the control file for any auxiliary files not required, for the "none" option for tty output, and for the "0" option for the user subroutine number. The code will always write an input check file and a restart file, so if names are not provided by the user, the defaults will be used. If an output file name is not specified, the generalized output is written to the terminal.

Input variable	Format	Opt/Req	Default	Description
nmfil(1)	character*100	Opt	fehmn.files	Control file name (this line is not included in the control file).
nmfil( 2)	character*100	Req	fehmn.dat	Main input file name.
nmfil( 3)	character*100	Opt	not used	Geometry-data input file name.
nmfil( 4)	character*100	Opt	not used	Zone-data input file name.
nmfil( 5)	character*100	Opt	terminal	Main output file name.
nmfil( 6)	character*100	Opt	not used	Restart input file name.
nmfil( 7)	character*100	Opt	fehmn.fin	Restart output file name.
nmfil( 8)	character*100	Opt	not used	Simulation-history output file name.
nmfil( 9)	character*100	Opt	not used	Solute-history output file name.
nmfil(10)	character*100	Opt	not used	Contour-plot output file name (required if using avs option in cont macro).
nmfil(11)	character*100	Opt	not used	Dual-porosity or double-porosity/double-permeability contour-plot output file name.
nmfil(12)	character*100	Opt	not used	Coefficient-storage output file name.
nmfil(13)	character*100	Opt	fehmn.chk	Input-check output file name.
tty_flag	character*4	Opt	none	Terminal output flag: all, some, none.
usub_num	integer	Opt	0	User subroutine call number.

The following are examples of input control files.

tape5.dat	
tape5.dat	
tape5.dat	
tape5.out	
tape5.his	
tape5.trc	
tape5.con	
tape5.chk	
some	
0	

/groupdir/c14-3
/groupdir/grid-402
/groupdir/c14-3
c14-3.out
/groupdir/c14-3.ini
c14-3.fin
c14-3.his
c14-3.trc
c14-3.con
c14-3.dp
c14-3.stor
c14-3.chk
none
0

## 6.2.2 Optional input files

The data for any of the FEHM macros (with the exception of **coor** and **elem**) may be entered in an alternate input file. To use this option, the keyword "file" must appear on the input line immediately following the control statement (macro name). The line immediately following this keyword will contain the name of the alternate input file. The contents of the alternate input file consist of the regular full macro description: the macro name followed by the data. The entries in the optional input file may be preceded or followed by comments using the "#" designator (see discussion on page 19). As with regular macro input, comments may not be embedded within the data block.

Group 1 - LOCKEYWORD

Group 2 - LOCFILENAME

Input variable	Format	Description
LOCKEYWORD	character*4	Keyword "file" to designate an auxiliary input file is used.
LOCFILENAME	character*100	Name of the optional data input file.

The following illustrate the use of an optional file and its contents.

rock
file
rockfile

File "rockfile":

# Auxiliary file used for rock macro input					
rock					
1	140	1	2563.	1010.	0.3500
# End of rock macro input					

### 6.2.3 Control statement **adif** (optional)

Air-water vapor diffusion.

Group 1- TORT

Input variable	Format	Description
TORT	real	Tortuosity for air-water vapor diffusion.

### 6.2.4 Control statement **airwater** (optional)

Isothermal air-water two-phase simulation.

Several macros are affected if the air module is enabled. These are:

- pres** - Because the air-water formulation is two-phase at all times, care should be taken to insure that IEOSD is always specified to be 2. Likewise, saturations (not temperatures) are used.
- init** - This macro should not be used because the saturation values cannot be specified.
- flow** - A variety of different flow and boundary values are input with this macro when the macro **airwater** is also used. See description of control statement **flow** (page 37).

Group 1 - ICO2D

Group 2 - TREF, PREF

Input variable	Format	Description
ICO2D	integer	Determines the type of air module used. ICO2D = 1, 1-degree-of-freedom solution to the saturated-unsaturated problem is produced. This formulation is similar to the Richard's Equation. ICO2D = 2, 1-degree-of-freedom solution is obtained assuming only gas flow with no liquid present. ICO2D = 3, full 2-degrees-of-freedom solution. All other values are ignored. The default is [3].
TREF	real	Reference temperature for properties (°C).
PREF	real	Reference pressure for properties (MPa).



The following is an example of **airwater**:

airwater
3
20. 0.1

### 6.2.5 Control statement **alti** (optional)

Alternate element and coordinate input. Not supported in this version.

Group 1 - CC, N

Group 2 - INFL

Input variable	Format	Description
CC	character*4	Input file type (ment: mentat mesh generator; ptrn: patran mesh generator).
N	integer	Number of nodes in the grid.
INFL	character*100	Name of alternate element- and coordinate-data input file.

### 6.2.6 Control Statement **boun** (either **boun** or **flow** is required)

Implement boundary conditions and sources or sinks. Input may be time dependent and cyclic. Time-step sizes may also be adjusted.

Group 1 - KEYWORD

Group 2 - NTIMES, (TIME\_MODEL(I), I=1, NTIMES) (KEYWORDS 'ti' and 'cy') or (VARIABLE\_MODEL(I), I=1, NTIMES) (repeated as needed)

Group 3 - JA, JB, JC, MODEL\_NUMBER (JA, JB, JC - defined on page 19)

Input variable	Format	Description
KEYWORD	character*4	<p>Keyword specifying a model designation, time for boundary condition or source changes, or actual variable or source change. Keywords are:</p> <ul style="list-style-type: none"> <li>model - new model definition to follow</li> <li>ti - time sequence for changes to follow</li> <li>cy - cyclic time sequence for changes to follow</li> <li>sa - air-source sequence for changes to follow</li> <li>sw - water-source sequence for changes to follow</li> <li>se - enthalpy-source sequence for changes to follow</li> <li>dsa - distributed air-source sequence for changes to follow</li> <li>dsw - distributed water-source sequence for changes to follow</li> <li>dse - distributed enthalpy-source sequence for changes to follow</li> <li>s - fixed saturation sequence for changes to follow</li> <li>hd - fixed hydraulic-head sequence for changes to follow</li> <li>pw - fixed water-pressure sequence for changes to follow</li> <li>pa - fixed air-pressure sequence for changes to follow</li> <li>pwo - fixed water-pressure sequence for changes to follow (constrained to outflow only)</li> <li>pao - fixed air-pressure sequence for changes to follow (constrained to outflow only)</li> <li>en - fixed enthalpy sequence for changes to follow</li> <li>t - fixed temperature sequence for changes to follow</li> <li>h - fixed humidity sequence for changes to follow (must be used with van Genuchten relative-permeability model)</li> <li>ft - fixed flowing-temperature sequence for change to follow. By flowing temperature, we mean the temperature of the inflow stream for a specified source. If no influx source occurs where this condition is applied, it will be ignored.</li> <li>if - impedance factor for use with fixed water-pressure boundary condition. If left out, the impedance factor will be set to the volume of the grid cell.</li> <li>ts - time-step sequence for changes to follow</li> <li>end - signifies end of keyword input; a blank line will also work.</li> </ul> <p>NOTE: Either KEYWORD 'ti' or 'cy' must be the first KEYWORD after a new model (KEYWORD 'model') is started.</p>
NTIMES	integer	<p>Number of time changes for boundary condition or source specification. This parameter is only associated with KEYWORDS 'ti' and 'cy', i.e., those that are associated with time sequences.</p>
TIME_MODEL	real	<p>Times for changes in boundary conditions or sources.</p>

Input variable	Format	Description
VARIABLE_MODEL	real	New values for boundary conditions or sources.
MODEL_NUMBER	integer	Number referring to the numerical order in which the models (begin with KEYWORD 'model') were input.

NOTE: The keywords concerning time 'ti' and 'cy' require additional details. The time must start at 0.0. This provides the initial boundary and source information. If the inputs for 'ti' or 'cy' do not start at 0.0, the time is added with the conditions at 0.0 set to 0.0. The 'cy' keyword involves a cyclic changing of conditions. In our procedure, the cycle ends at the last specified time. Thus, the code reverts to the first specified time values. Because of this, the boundary conditions and sources for the last time changes are always set to the first time values.

The following is an example of **boun**:

```

boun
model 1
  cy
  4 0.0 1.e1 1.e2 1.e5
  sw
  -1.e-4 -1.e-5 -1.e-3 -1.e-4
  ft
  20.0 50.0 50.0 20.0
model 2
  ti
  2 0.0 1.e20
  pw
  0.1 0.1
  ft
  20.0 20.0

26 26 1 1
27 27 1 2

```

In this example two models are defined. The first model cyclically changes the water source in a 1.e05-day cycle. Also, in model 1, the flowing temperature was alternated between 20°C and 50°C. Note that the water source at 1.e05 days equals that at 0.0 days. The second model uses a time change that occurs at 1.e20 days. This late change effectively removes any time variance from Model 2. Model 2 also has a fixed water-pressure condition. The models are applied to nodes 26 and 27 in the last two lines. The 'cy' keyword entries show that the time cycle ends at 1.e05 days, at which time, the cycle reverts to 0.0 days.

### 6.2.7 Control statement **bous** (optional)

Constant density and viscosity for the flow terms (Bousinesq approximation). NOTE: where the **bous** macro is used, the gravity term in the air phase is set to zero.

Group 1 - IBOUS

Input variable	Format	Description
IBOUS	integer	Parameter that enables the <b>bous</b> macro IBOUS = 1 enabled. IBOUS $\neq$ 1 disabled (default).

**6.2.8 Control statement cap** (no longer used, see macro **rlp**, page 55)

**6.2.9 Control statement cond** (required for nonisothermal problem)

Assign thermal conductivities of the rock.

Group 1 - JA, JB, JC, THXD, THYD, THZD (JA, JB, JC - defined on page 19)

Input variable	Format	Default	Description
THXD	real	1.e-30	Thermal conductivity in the x-direction ( $\frac{W}{m \cdot K}$ ).
THYD	real	1.e-30	Thermal conductivity in the y-direction ( $\frac{W}{m \cdot K}$ ).
THZD	real	1.e-30	Thermal conductivity in the z-direction ( $\frac{W}{m \cdot K}$ ).

The following is an example of **cond**:

cond					
1	140	1	1.00e-00	1.00e-00	0.00e-00

**6.2.10 Control statement cont** (optional)

Contour-data output format, output time-step intervals, and time intervals.

Group 1 - NCNTR, CONTIM

An alternative form of input for macro **cont** is possible. This is:

Group 1 - ALTC, NCNTR, CONTIM

Group 2 - CHDUM (only input if ALTC is 'avs')

FEHM will automatically distinguish between the alternative input formats. When keywords are used, they must be entered starting in the first column. The contour data will be output whenever either of the interval criteria are satisfied.

For AVS output, if the *material* keyword is selected, the following material property values will be written for each node: permeability in the x-, y-, and z-directions, thermal conductivity in the x-, y-, and z-directions, porosity, rock specific heat, capillary pressure, relative-permeability model being used, and capillary-pressure model being used. If *vapor* or *liquid* are selected, *pressure* or *velocity* must also be defined (otherwise, no data for

Input variable	Format	Description
ALTC	character*4	Keyword specifying the type of contour output wanted (avs, fehm, free, ment, ptrn): 'avs' produces contour plot files compatible with the AVS postprocessor. 'fehm' produces a binary output file. The same contour plot file is produced using the first form of Group1 input. 'free' produces a free-format contour plot file. 'ment' produces a contour plot file compatible with the MENTAT postprocessor. 'ptrn' produces a contour plot file compatible with the PATRAN postprocessor.
NCNTR	integer	<i>time-step</i> interval for contour plots (number of time steps). Output contour information each NCNTR time steps.
CONTIM	real	<i>Time</i> interval for contour plots (days). In addition to output each NCNTR time steps, output contour information each CONTIM days.
CHDUM	character*72	Keyword specifying type of AVS contour-plot data files to be created in AVS UCD format, either formatted (ASCII) or unformatted (binary). Keywords are entered one per line and terminated with 'endavs'. Valid keywords (case insensitive) are: (m)aterial - output contour values for material properties. (l)iquid - output contour values for liquid phase. (va)por - output contour values for vapor phase. (ve)locity - output velocity values. (dp)dp - output contour values for dual-permeability nodes. (p)ressure - output pressure values. (t)emperature - output temperature values. (s)aturation - output saturation values. (c)oncentration - output solute-concentration values. (f)ormatted - output data in ASCII format. (u)nformatted - output data in binary format. (e)ndavs - last keyword entered. If a format keyword is not entered, the default is 'formatted'. The default for data keywords is 'off.' The letters given in ( ) are sufficient to identify the keyword.

these values will be written). *velocity* will result in vector values; *pressure* values will be scalar. If *concentration* is selected, values will be output only if NSPECI is defined for tracer solutions. See the control statement **trac** for a description of NSPECI for solutes.

The following are examples of **cont**. For the first example, FEHM binary-format contour output files will be written every 100 time steps and for each 1.e20 days. The second example invokes AVS contour output. AVS UCD binary files will be written for every 100 time steps and 1.e20 days. The resulting files will include a log file, geometry file, plus header and data files for the following: material properties, solute concentrations, liquid velocities, pressures, and temperatures.

```
cont
100      1.e20
```

```
cont
avs      100      1.e20
liquid
velocity
con
pressure
temp
mat
unformatted
end
```

### 6.2.11 Control statement **coor** (required)

Node coordinate data. These data are usually created by a mesh-generation program, then cut and copied into the input file or a separate geometry-data input file. The mesh must be a right-handed coordinate system.

Group 1 - N

Group 2 - MB, CORD1, CORD2, CORD3

To end the control section a blank line is entered.

Input variable	Format	Description
N	integer	Number of nodes in the grid.
MB	integer	Node number. If MB < 0, then the difference between the absolute value of MB and the previously read absolute value of MB is used to generate intermediate values by interpolation.
CORD1	real	X-coordinate of node MB (m).
CORD2	real	Y-coordinate of node MB (m).
CORD3	real	Z-coordinate of node MB (m).

The following is an example of **coor**:

```
coor
140
  1      0.00000      200.00000      0.00000
  2      12.50000      200.00000      0.00000
  .
  .
  .
 10      212.50000      200.00000      0.00000
  .
  .
  .
140      300.00000      0.00000      0.00000
```

## 6.2.12 Control statement ctrl (required)

Assigns various control parameters needed for equation-solver and matrix-solver routines.

Group 1 - MAXIT, EPM, NORTH

Group 2 - JA, JB, JC, IGAUS (JA, JB, JC - defined on page 19)

Group 3 - AS, GRAV, UPWGT

Group 4 - IAMM, AIAA, DAYMIN, DAYMAX

Group 5 - ICNL, LDA

Input variable	Format	Default	Description
MAXIT	integer		Maximum number of iterations allowed in either the overall Newton cycle or the inner cycle to solve for the corrections at each iteration. If MAXIT < 0, then the maximum number of iterations is ABS(MAXIT), but the minimum number of iterations is set to 2. [10]
EPM	real		Tolerance for Newton cycle (nonlinear equation tolerance). [1.e-5]
NORTH	integer		Number of orthogonalizations in the linear-equation solver. [8]
IGAUS	integer	1	The order of partial Gauss elimination (1 or 2 is recommended). Larger values increase memory use but may be necessary for convergence.
AS	real		Implicitness factor. [1] AS ≤ 1, use standard pure implicit formulation. AS > 1, use second-order implicit method.
GRAV	integer		Direction of gravity. GRAV = 0, no gravity is used. GRAV = 1, x-direction. GRAV = 2, y-direction. GRAV = 3, z-direction. A value for gravity of 9.81 m/s <sup>2</sup> is used in the code when GRAV ≠ 0. If GRAV > 3, GRAV is set equal to 3.
UPWGT	real		Value of upstream weighting (0.5 ≤ UPWGT ≤ 1.0). If UPWGT < 0.5, UPWGT is set to 0.5. If UPWGT > 1.0, UPWGT is set to 1.0.
IAMM	integer		Maximum number of iterations for which the code will multiply the time-step size. If this number of time steps is exceeded at any time, the time step will not be increased for the next time. [7-10]
AIAA	real	1	Time-step multiplier. [1.2-2.0]
DAYMIN	real		Minimum time-step size (days).
DAYMAX	real		Maximum time-step size (days).

Input variable	Format	Default	Description
ICNL	integer		Parameter that specifies the geometry. ICNL = 0, three-dimensional. ICNL = 1, x-y plane. ICNL = 2, x-z plane. ICNL = 3, y-z plane. ICNL = 4, x-y radial plane, (radius is x). ICNL = 5, x-z radial plane, (radius is x). ICNL = 6, y-z radial plane, (radius is y).
LDA	integer	0	Parameter that specifies the external storage of geometric coefficients. LDA = +1, element coefficients are read from file <i>filen.stor</i> , and no coefficients are calculated in the code. LDA = 0, element coefficients are calculated in the code and not saved. LDA = -1, element coefficients are calculated in the code and saved on file <i>filen.stor</i> . It should be noted that if the coefficients are read from a file (LDA = 1), then the macro <b>finv</b> is ignored as well as information read from macros <b>elem</b> and <b>coor</b> .

The following is an example of **ctrl**:

ctrl				
40	1.e-7	8		
1	140	1		1
1.0	0.0	1.0		
40	1.2	0.1		60.0
1	0			

### 6.2.13 Control statement **dof** (Not implemented)

### 6.2.14 Control statement **dpgp** (optional)

Double-porosity/double-permeability formulation. There are two sets of parameter values at any nodal position for which property values must be defined. Nodes 1 to N (see macro **coor**, page 29, for definition of N) represent the fracture nodes and nodes N + 1 to 2N, the matrix material. When zones are used with the **dpgp** macro, additional zones are automatically generated. See instructions for the macro **zone** for a more detailed description. The **dpgp** parameters are only defined for the first N nodes.

Group 1 - IDPDP

Group 2 - JA, JB, JC, VOLFD1 (JA, JB, JC - defined on page 19)

Group 3 - JA, JB, JC, APUV1 (JA, JB, JC - defined on page 19)

The volume fraction VOLFD1 is related to the total volume by

$$VOLFD1 + VOLFD2 = 1.0 ,$$



Input variable	Format	Default	Description
IDPDP	integer		Solution descriptor for double-porosity/double-permeability solution. IDPDP = 0, information is read but not used. IDPDP $\neq$ 0, <b>dpdp</b> solution is implemented.
VOLFD1	real	1.	Volume fraction for fracture node.
APUV1	real	10.	Length scale for matrix nodes (m).

where VOLFD2 is the volume fraction of the matrix node. If permeability model IRLP = 4 is selected in control statement **rlp**, VOLFD1 is calculated from RP15 (fracture porosity) in that control statement.

The following is an example of **dpdp**:

dpdp				
1				
1	140	1	0.005	
1	140	1	0.10	

### 6.2.15 Control statement dual (optional)

Dual-porosity formulation. There are three sets of parameter values at any nodal position for which property values must be defined. Nodes 1 to N (see macro **coor**, page 29, for definition of N) represent the fracture nodes, nodes N + 1 to 2N, the first matrix material, and nodes 2N + 1 to 3N, the second matrix material. When zones are used with the **dual** macro, additional zones are automatically generated (see instructions for the macro **zone**, page 74, for a more detailed description). The **dual** parameters are only defined for the first N nodes.

Group 1 - IDUALP

Group 2 - JA, JB, JC, VOLFD1 (JA, JB, JC - defined on page 19)

Group 3 - JA, JB, JC, VOLFD2 (JA, JB, JC - defined on page 19)

Group 4 - JA, JB, JC, APUVD (JA, JB, JC - defined on page 19)

Input variable	Format	Default	Description
IDUALP	integer		Solution descriptor for dual-porosity solution. IDUALP = 0, information is read but not used. IDUALP $\neq$ 0, dual-porosity solution is implemented.
VOLFD1	real	0.001	Volume fraction for fracture portion of the continuum.
VOLFD2	real	0.5	Volume fraction for the first matrix portion of the continuum.
APUVD	real	5.	Length scale for the matrix nodes (m).

The volume fractions VOLFD1 and VOLFD2 are related to the total volume by

$$VOLFD1 + VOLFD2 + VOLFD3 = 1.0 ,$$

where VOLFD3 is the volume fraction of the second matrix node. If permeability model IRLP = 4 is selected in control statement **rlp**, VOLFD1 is calculated from RP15 (fracture porosity) in that control statement.

The following is an example of **dual**:

dual				
1				
1	140	1		0.006711409
1	140	1		0.335570470
1	140	1		0.10

## 6.2.16 Control statement elem (required).

Element connectivity data. These data are created by a mesh-generation program, then cut and copied into the input file or a separate geometry-data input file.

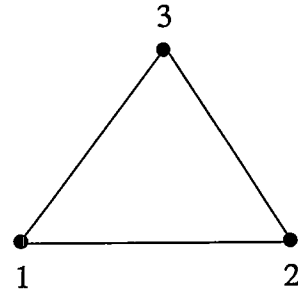
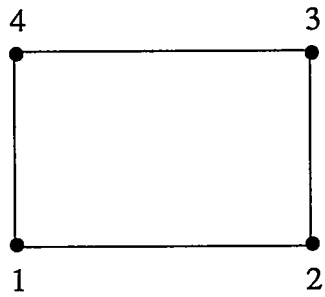
Group 1 - NS, NEI

Group 2 - MB, NELM (1), NELM (2), . . . , NELM (NS)

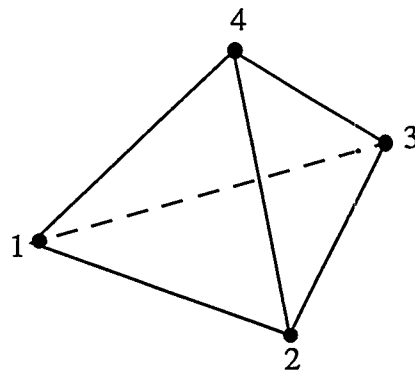
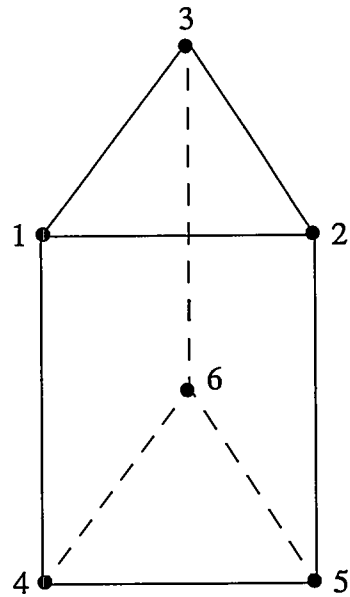
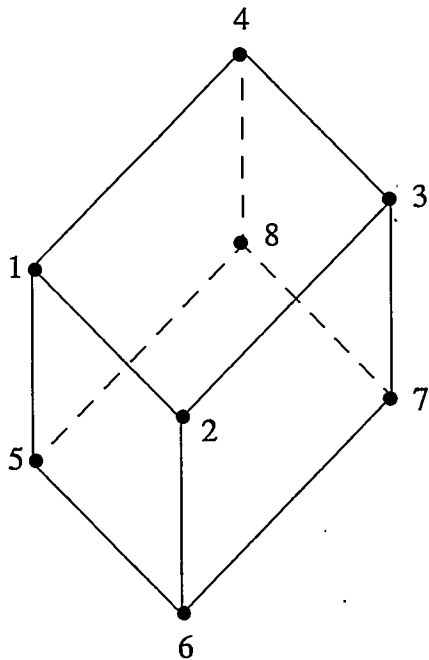
IF  $NS < 0$ , then ABS(NS) is interpreted as the number of nodes per element.  $NS < 0$  signals the code to make rectangles (or bricks in three dimensions) a sum of triangles (or tetrahedrons). This approach provides more stability in nonlinear problems with a distorted mesh. Figure 2 shows available element types and the nodal numbering convention. To end the control section a blank line is entered.

Input variable	Format	Description
NS	integer	Number of nodes per element.
NEI	integer	Number of elements.
MB	integer	Element number. If $MB < 0$ , then the difference between the absolute value of MB and the previous absolute value of MB is used to generate intermediate values by interpolation in the code.
NELM (1)	integer	First node of element MB.
NELM (2)	integer	Second node of element MB.
⋮	⋮	⋮
NELM (NS)	integer	Last node of element MB.

## 2-D



## 3-D



**Figure 2. Elements available with FEHM in 2-D and 3-D problems showing nodal numbering convention.**

The following is an example of **elem**:

elem				
4	117			
1	15	16	2	1
2	16	17	3	2
.	.	.	.	.
.	.	.	.	.
.	.	.	.	.
10	24	25	11	10
11	25	26	12	11
12	26	27	13	12
.	.	.	.	.
.	.	.	.	.
.	.	.	.	.
116	138	139	125	124
117	139	140	126	125

### 6.2.17 Control statement eos (optional)

Equation of State. Provide the code with alternate thermodynamic properties for the liquid and/or vapor phases. (This is one way in which the code may be instructed to simulate nonisothermal, single-phase air. It may also be used to make comparisons between the code and analytical solutions that use different equations of state.)

Group 1 - IIEOSD, IPSAT, ITSAT

Group 2 - EW1, EW2, EW3, EW4, EW5, EW6, EW7, EW8, EW9, EW10, EW11

Group 3 - EV1, EV2, EV3, EV4, EV5, EV6, EV7, EV8, EV9, EV10, EV11

For the calculation of vapor density and its derivatives, the ideal gas law is used instead of a linear relationship. Thus, EV4 and EV5 are not used but are included so the format is the same as that for the liquid parameters in Group 2.

Input variable	Format	Description
IIEOSD	integer	Equation-of-state reference number. When IIEOSD = 1 or 2 are used, they refer to the high- and low-pressure data sets, respectively, in FEHM. For these values, the input in Group 2 and Group 3 will be ignored after it is entered. When any value other than 1 or 2 are used, the user-defined equation of state is used with Groups 2 and 3 for input.
IPSAT	integer	Parameter to set vapor pressure to zero. If IPSAT $\neq$ 0, the vapor pressure is set to zero, otherwise the vapor pressure is calculated in the code.
ITSAT	integer	Parameter to adjust the saturation temperature. If ITSAT < 0, the saturation temperature is set to -1000°C. If ITSAT > 0, the saturation temperature is set to 1000°C. If ITSAT = 0, the calculated value is used.
EW1	real	Liquid reference pressure (MPa).
EW2	real	Liquid reference temperature (°C).

Input variable	Format	Description
EW3	real	Liquid reference density (kg/m <sup>3</sup> ).
EW4	real	Derivative of liquid density with respect to pressure at reference conditions.
EW5	real	Derivative of liquid density with respect to temperature at reference conditions.
EW6	real	Liquid reference enthalpy (MJ/kg).
EW7	real	Derivative of liquid enthalpy with respect to pressure at reference conditions.
EW8	real	Derivative of liquid enthalpy with respect to temperature at reference conditions.
EW9	real	Liquid reference viscosity.
EW10	real	Derivative of liquid viscosity with respect to pressure at reference conditions.
EW11	real	Derivative of liquid viscosity with respect to temperature at reference conditions.
EV1	real	Vapor reference pressure (MPa).
EV2	real	Vapor reference temperature (°C).
EV3	real	Vapor reference density (kg/m <sup>3</sup> ).
EV4	real	Not used, included only to maintain a similar format to Group 2.
EV5	real	Not used, included only to maintain a similar format to Group 2.
EV6	real	Vapor reference enthalpy (MJ/kg).
EV7	real	Derivative of vapor enthalpy with respect to pressure at reference conditions.
EV8	real	Derivative of vapor enthalpy with respect to temperature at reference conditions.
EV9	real	Vapor reference viscosity.
EV10	real	Derivative of vapor viscosity with respect to pressure at reference conditions.
EV11	real	Derivative of vapor viscosity with respect to temperature at reference conditions.

### 6.2.18 Control statement exrl (optional)

Allows the user to choose explicit relative permeability.

Group 1 - IEXRLP

Input Variable	Format	Description
IEXRLP	integer	If IEXRLP=1, then explicit relative permeability. Otherwise not enabled.

### 6.2.19 Control statement finv (optional)

No input is associated with this macro. When invoked, the code will perform finite-volume calculations instead of finite-element calculations for flow terms—this may improve accuracy on nonorthogonal grid systems. Anisotropic properties (permeability, conductivity) are not supported with this macro. In this case, the values for permeability in the x-direction from control statement **perm** are used.

### 6.2.20 Control statement flow (required for flow problem)

Flow data. Source and sink parameters are input and may be used to apply boundary conditions. Note that the alternative definitions (isothermal conditions) apply when control statement **airwater** is used.

Group 1 - JA, JB, JC, SKD, EFLOW, AIPED (JA, JB, JC - defined on page 19)

If the porosity of the node is zero, then there is only a temperature solution, and the code forms a source proportional to the enthalpy difference. The source term is given by  $Q = AIPED \cdot (E - EFLOW)$ , where E is the in-place enthalpy and EFLOW is a specified enthalpy.

Input variable	Format	Default	Description
<b>Nonisothermal</b>			
SKD	real	0.	Heat and mass source strength (kg/s), heat only (MJ/s). Negative value indicates injection into the rock mass.
EFLOW	real	0.	Enthalpy of fluid injected (MJ/kg). If the fluid is flowing from the reservoir, then the in-place enthalpy is used. If EFLOW < 0, then ABS(EFLOW) is interpreted as a temperature (°C) and the enthalpy (assuming water only) calculated accordingly. In heat-only problems with EFLOW < 0, the node is in contact with a large heat pipe that supplies heat to the node through an impedance AIPED so as to maintain its temperature near ABS (EFLOW). Large values (approximately 1000) of AIPED are recommended.
AIPED	real	0.	Impedance parameter. If AIPED is nonzero, the code interprets SKD as a flowing wellbore pressure (MPa) with an impedance ABS(AIPED). If AIPED < 0, flow is only allowed out of the well. For heat only, AIPED is the thermal resistance. If AIPED = 0, SKD is flow rate. If AIPED ≠ 0 and SKD = 0, the initial value of pressure will be used for the flowing pressure.

Input variable	Format	Default	Description
<b>Isothermal air-water</b>			
Case 1: AIPED = 0 (constant mass rate, 1- or 2-phase source or sink)			
SKD	real	0.	Mass source strength (kg/s). Negative value indicates injection into the rock mass.
EFLOW	real	0.	<p>a) <math>EFLOW \geq 0</math>, EFLOW is source liquid saturation.  <math>Q_w = SKD \cdot EFLOW</math> (kg/s).  <math>Q_a = SKD \cdot (1 - EFLOW)</math> (kg/s).</p> <p>b) <math>EFLOW &lt; 0</math>, ABS(EFLOW) is the source air pressure (MPa).  <math>Q_w = SKD</math> (kg/s).  <math>Q_a = 1.0 \cdot (P_a - ABS(EFLOW))</math> (kg/s).</p> <p>In the above and following relations, <math>Q_w</math> is the source term for water, <math>Q_a</math> is the source term for air, and <math>P_a</math> is the in-place air pressure. The second case works well in situations in which inflow is specified and it is desired to hold the air pressure at a constant value.</p>
Case 2: AIPED > 0 (constant pressure, constant liquid saturation source or sink)			
SKD	real	0.	Specified source air pressure (MPa).
EFLOW	real	0.	<p>a) <math>EFLOW &lt; 0</math>, air only source.  <math>Q_a = AIPED \cdot (P_a - SKD)</math> (kg/s).</p> <p>b) <math>0 &lt; EFLOW \leq 1</math>, EFLOW is specified source liquid saturation. For <math>SKD \geq 0</math>, 2-phase source.  <math>Q_a = AIPED \cdot (P_a - SKD)</math> (kg/s).  <math>Q_w = AIPED \cdot (S_l - EFLOW)</math> (kg/s).</p> <p>For <math>SKD &lt; 0</math>, water only source. <math>Q_a = 0</math>.</p> <p>In the above relation, <math>S_l</math> is the in-place liquid saturation.</p>
AIPED	real		Impedance parameter. A large value is recommended ( $10^2 - 10^6$ ) to create a flow term large enough to maintain constant pressure.
Case 3: AIPED < 0 (Outflow only , if $P_l > SKD$ )			
SKD	real	0.	Pressure above which outflow occurs (MPa).
EFLOW	real	0.	Not used.

Input variable	Format	Default	Description
AIPED	real	0.	Impedance parameter. $Q_w = ABS(AIPED) \cdot R_l / \mu_l (P_l - SKD) \text{ (kg/s)}$ where $R_l$ is the water relative permeability and $\mu_l$ is the water viscosity.

The following is an example of **flow**:

flow					
88	88	1	0.050	-25.0	0.
14	14	1	3.600	-160.0	1.

### 6.2.21 Control statement flo2 (optional)

Group 1 - JA, JB, JC, JD, SKD, EFLOW, AIPED (SKD, EFLOW, AIPED - defined on page 37 under control statement **flow**)

Multiple lines of input may be used terminated by a blank line.

Input variable	Format	Description
JA	integer	Indices used to define planes in a 3-D simulation with a regular numbering pattern. The flow rates are defined within the inner loop of the do loops: DO JK = JA, JB KL = JK - JA DO IJ = JA + KL, JC + KL, KD ... ENDDO ENDDO
JB	integer	
JC	integer	
JD	integer	

### 6.2.22 Control statement flxo (optional)

Mass flux between two nodes is output by choosing this control statement.

Group 1 - NFLX

Group 2 - IFLX1, IFLX2 (repeated NFLX times)

Group 3 - X1, Y1, Z1 (as needed)

Group 4 - X2, Y2, Z2 (as needed)

Input variable	Format	Description
NFLX	integer	Number of internode fluxes to be calculated.
IFLX1	integer	First node to be used in flux calculation.
IFLX2	integer	Second node to be used in flux calculation. If IFLX2 = 0, then the node connected to IFLX1 with the greatest internodal distance is used to calculate the mass flux.



Input variable	Format	Description
X1	real	Coordinates of the first node to be used in flux calculation. Used only for those nodes for which IFLX1 < 0.
Y1	real	
Z1	real	
X2	real	Coordinates of the second node to be used in flux calculation. Used only for those nodes for which IFLX2 < 0.
Y2	real	
Z2	real	

If IFLX1 < 0, then after all IFLX1 and IFLX2 values are read, coordinates X1, Y1, and Z1 are read, and the node nearest to these coordinates is used. If IFLX2 < 0, coordinates for the second node are read in on another line. The code cycles through each IFLX1 and IFLX2 in this manner, reading coordinates when needed. Results are written to the screen, if tty output is enabled, and to the output file **iout**.

### 6.2.23 Control Statement head (optional)

No input is associated with this control statement. Uses hydraulic head values as input and output. Enables **bous** macro (Bousinesq approximation) automatically. Enables **airw** macro (air-water isothermal) automatically. It affects the **pres** and **flow** macros by requiring head information where pressure values were previously required.

### 6.2.24 Control statement hflx (optional)

Group 1 - JA, JB, JC, QFLUX, QFLXM (JA, JB, JC - defined on page 19)

A negative heat flux indicates heat flow into the reservoir.

Input variable	Format	Default	Description
QFLUX	real	0.	If QFLXM = 0, then QFLUX is the heat flux (MW). If QFLXM ≠ 0, then QFLUX is a temperature (°C) and the heat flux is calculated according to the formula: $Q_H = QFLXM(TL - QFLUX) \text{ (MW).}$
QFLXM	real	0.	If QFLXM ≠ 0, multiplier for heat flux equation given in QFLUX description (MW/°C). This must be large for large temperature gradients or when a constant temperature must be maintained.

The following is an example of **hflx**:

hflx				
401	410	1	-0.001	0.0

### 6.2.25 Control statement ice (optional)

Ice-phase calculations (not tested).

Group 1 - ICE, SIIN, TMELT

Group 2 - JA, JB, JC, SII (JA, JB, JC - defined on page 19)

Input variable	Format	Description
ICE	integer	Solution descriptor for ice solution. ICE = 0, information is read but not used. ICE ≠ 0, ice solution is implemented.
SIIN	real	Default value for ice saturation (used when ice saturation SII in Group 2 is set to 0 at any node).
TMELT	real	Freezing temperature of water (°C).
SII	real	Ice saturation. The default value is [0].

### 6.2.26 Control statement init (required if macro pres not used)

Set initial pressure and temperature at all nodes.

Group 1 - PEIN, TIN, TIN1, GRAD1, DEPTH, TIN2, GRAD2, QUAD

Note that the macro **pres** may overwrite some of the values that are set by macro **init**.

Input variable	Format	Description
PEIN	real	Initial value of pressure (MPa). If initial values are read from the read file (iread), then this value is ignored. If gravity is present, this is the value of the pressure at node 1, and the other nodal pressures are adjusted by applying the hydraulic head. Absolute pressures are used. Pressure as a function of depth is calculated with $TIN < 0$ .
TIN	real	Initial value of temperature (°C). If $TIN \leq 0$ , then the initial temperatures are calculated using the temperature-gradient formulas given below.
TIN1	real	Defined in formulas below (°C).
GRAD1	real	Defined in formulas below (°C/m).
DEPTH	real	Defined in formulas below (m).
TIN2	real	Defined in formulas below (°C).
GRAD2	real	Defined in formulas below (°C/m).
QUAD	real	Defined in formulas below (°C/m <sup>2</sup> ).
$T = TIN1 + GRAD1 \times Z \quad 0 \leq Z \leq DEPTH$		
$T = TIN2 + GRAD2 \times Z + QUAD \times Z^2 \quad Z > DEPTH$		

The following is an example of **init**:

init	3.6	0.0	240.	0.	0.	240.	0.	0.
------	-----	-----	------	----	----	------	----	----

### 6.2.27 Control statement **iter** (optional)

If the user is not familiar with the linear-equation-solver routines in FEHM, control statement **iter** should not be used.

Group 1 - G1, G2, G3, TMCH, OVERF

Group 2 - IRDOF, ISLORD, IBACK, ICOUPL, RNMAX

Input variable	Format	Default	Description
G1	real	1.e-6	Multiplier for the linear-convergence region of the Newton-Raphson iteration.
G2	real	1.e-6	Multiplier for the quadratic-convergence region of the Newton-Raphson iteration.
G3	real	1.e-3	Tolerance for the adaptive implicit method (multiplying factor for Newton-Raphson tolerance).
TMCH	real	1.e-9	Machine tolerance. If satisfied by the residual norm, the Newton iteration is assumed to be complete. If TMCH is < 0, the ABS(TMCH) is used as a tolerance for each equation at each node. Convergence is achieved if the residual of every equation at every node is < ABS(TMCH).
OVERF	real	1.1	Over-relaxation factor for passive nodes in adaptive implicit method.
IRDOF	integer	0	Enables the reduced degree-of-freedom method. [0] Set to 0 if reduced degrees of freedom are not required. When IRDOF = 1, a reduced degree of freedom from 3 to 2 or 3 to 1 is used. When IRDOF = 2, a reduced degree of freedom from 3 to 2 is used. If IRDOF = 11, then an air-only solution is found for the isothermal air-water process model. If IRDOF = -1, then the residual for the air equation with <b>airw</b> macro is ignored. If IRDOF = 14, then the saturated-unsaturated 1-degree-of-freedom solution scheme is used (macro <b>airw</b> only).

Input variable	Format	Default	Description																												
ISLORD	integer	0	Reordering parameter. The ordering can be understood by labeling the mass equation as 1, the heat equation as 2, and the noncondensable gas equation (if it exists) as 3. The value of ISLORD and the corresponding equation order is given below. The ordering has an effect on the speed of convergence of several solution algorithms but will not affect most users.																												
<table border="1"> <thead> <tr> <th>ISLORD</th><th>2 Degrees of Freedom</th><th>3 Degrees of Freedom</th><th>4 Degrees of Freedom</th></tr> </thead> <tbody> <tr> <td>0</td><td>1, 2</td><td>1, 2, 3</td><td>1, 2, 3, 4</td></tr> <tr> <td>1</td><td>2, 1</td><td>1, 3, 2</td><td>1, 3, 2, 4</td></tr> <tr> <td>2</td><td></td><td>2, 1, 3</td><td></td></tr> <tr> <td>3</td><td></td><td>2, 3, 1</td><td></td></tr> <tr> <td>4</td><td></td><td>3, 1, 2</td><td></td></tr> <tr> <td>5</td><td></td><td>3, 2, 1</td><td></td></tr> </tbody> </table>				ISLORD	2 Degrees of Freedom	3 Degrees of Freedom	4 Degrees of Freedom	0	1, 2	1, 2, 3	1, 2, 3, 4	1	2, 1	1, 3, 2	1, 3, 2, 4	2		2, 1, 3		3		2, 3, 1		4		3, 1, 2		5		3, 2, 1	
ISLORD	2 Degrees of Freedom	3 Degrees of Freedom	4 Degrees of Freedom																												
0	1, 2	1, 2, 3	1, 2, 3, 4																												
1	2, 1	1, 3, 2	1, 3, 2, 4																												
2		2, 1, 3																													
3		2, 3, 1																													
4		3, 1, 2																													
5		3, 2, 1																													
IBACK	integer		IRDOF parameter. [0] If IBACK = 1, SOR iterations are performed before call to solver. If IBACK = 2, SOR iterations are performed before call to SOLVER, and SOLVER is called twice.																												
ICOUPL	integer		Number of SOR iterations used in reduced degree of freedom methods. [0]																												
RNMAX	real		Maximum running time for problem before the solution is stopped (cpu minutes) (very large if not set with control statement <b>iter</b> ).																												

The following is an example of **iter**:

<b>iter</b>				
1.e-5	1.e-5	1.e-5	1.e-9	1.2
1	0	0	2	200.0

### 6.2.28 Control statement **itup** (optional)

Group 1 - IAD\_UP

Input variable	Format	Default	Description
IAD_UP	integer	100	Number of iterations after which the upwind directions are held constant. [2]

### 6.2.29 Control statement **iupk** (optional)

No input is associated with this control statement. If enabled, the full transmissibility term will be upwinded (including the intrinsic permeability). Otherwise, the fluid and relative-permeability part of the

transmissibility will be upwinded and the intrinsic permeability will be harmonically averaged.

### 6.2.30 Control statement ivfc (optional)

Enables volume-control subroutine. Not supported in this version.

### 6.2.31 Control statement mdnode (optional)

Enables extra connections to be made to nodes, which is useful for simulating wellbore connections, faults, and flow across internal boundaries.

Group 1 - NUM\_MD, MAX\_CON, IELIM, SX\_MULT

Group 2 - NODE, IPAR, NPAR (repeated NUM\_MD times)

Input variable	Format	Default	Description
NUM_MD	integer	0	Number of new connections to be entered.
MAX_CON	integer	0	Maximum number of new connections to a given node. This number does not include old connections. Thus, if a node was already connected to 5 neighboring nodes and two new connections were added to this node in this macro statement and this was the maximum number of connections added in this macro statement, then MAX_CON = 2.
IELIM	integer	0	IF IELIM $\geq$ 0, then no action. IF IELIM < 0, then nodal connections are eliminated as needed if redundant.
SX_MULT	real*8	1.0	Multiplier for equilibrium conditions.
NODE	integer	0	Node to which new connection is established.
IPAR	integer	0	IPAR is not used at present. Its value is ignored. However the entered number must be an integer.
NPAR	integer	0	NPAR is the new connected node. If NPAR = NODE, no new connection is established.

The following are examples of **mdnode**:

mdnode		
3	2	0 10
10	0	15
100	0	106
10	0	320

mdnode		
4	3	0 100
1	0	16
2	0	1
4	0	1
10	0	203

### 6.2.32 Control statement ngas (optional)

Noncondensable gas transport.

Group 1 - ICO2D

Group 2 - JA, JB, JC, PCO2 (JA, JB, JC - defined on page 19)

Group 3 - JA, JB, JC, CPNK (JA, JB, JC - defined on page 19)

Group 4 - JA, JB, JC, QCD (JA, JB, JC - defined on page 19)

Input variable	Format	Default	Description
ICO2D	integer	3	Solution descriptor for noncondensable gas transport. ICO2D = 1, the 3-degree-of-freedom solution will be reduced to a 1-degree-of-freedom problem. (See macro <b>iter</b> , the parameter ICOUPL is also set to 5 if ICO2D = 1.) ICO2D = 2, the 3-degree-of-freedom solution will be reduced to a 2-degree-of-freedom problem. (See macro <b>iter</b> , the parameter ICOUPL is also set to 5 if ICO2D = 2.) ICO2D = 3, full 3 degrees of freedom.
PCO2	real	0.	Initial partial pressure of noncondensable gas. If PCO2 < 0, then ABS (PCO2) is interpreted as a temperature and the partial pressure of the noncondensable gas is calculated according to the formula: $PCO2 = P_T - P_{SAT}(T)$ , where $P_T$ is the total pressure and $P_{SAT}(T)$ is the water-saturation pressure and is a function of temperature only.
CPNK	real	0.	If CPNK < 0, then ABS (CPNK) is the specified noncondensable pressure and will be held at that value. If CPNK > 0, then CPNK is the specified relative humidity and the saturation, $S_l$ , is calculated using the vapor-pressure lowering formula and the capillary-pressure formula: $Pcap(S_l) = \ln(h)\rho_l RT$ , where $Pcap$ is the capillary function, $h$ is the humidity, $R$ is the gas constant, $T$ is the temperature, and $\rho_l$ is the liquid density. Once the formula is solved, $S_l$ is held constant. The humidity condition is only enabled for van Genuchten capillary-function model. See macro <b>rlp</b> .
QCD	real	0.	Specified air flow rate (kg/sec).

The following is an example of **ngas**:

ngas				
3				
1	800	1	-20	
1	800	1	0.	
1	800	1	0.	

### 6.2.33 Control statement **nod2** (optional)

Specify the node numbers for which detailed file output is desired and alternate nodes for terminal output.

Group 1 - M, M2

Group 2 - MN (1), MN (2), . . . , MN (M)

Group 3 - X, Y, Z (as needed)

Group 4 - MNI(1), MNI(2), . . . , MNI(M2)

Group 5 - X, Y, Z (as needed)

Input variable	Format	Description
M	integer	Number of nodes for which information will be printed on the output file (iout ). If $M \leq 0$ , pressure and temperature will be written on the output file for all nodes, but no nodal parameter values will be printed in the history plot files. Group 2 is omitted if $M \leq 0$ .
M2	integer	Number of nodes for short list (terminal printout). If $M2 \leq 0$ , Group 4 is omitted.
MN	integer	M node numbers for which information will be printed on the output file (iout ). If a $MN(I) < 0$ , then coordinates are used to define that print-out node, and the coordinate sets (X, Y, Z) for each $MN(I) < 0$ are added after Group 2.
MNI	integer	M2 node numbers for which information will be printed on the terminal (short list). This group exists only if $M2 \neq 0$ . If $MNI(I) < 0$ , then coordinates are used to define the terminal output nodes, and the coordinate sets (X, Y, Z) for each $MNI(I) < 0$ are added after Group 4.
X	real	Coordinates of node for which information will be printed. One line for each MN or MNI $< 0$ . The code finds the node closest to the coordinate given. For 2-D problems, set $Z = 0$ . No input if no MN or MNI $< 0$ .
Y	real	
Z	real	

The following are examples of **nod2**:

nod2	
2	1
50	88
50	

nod2		
2	1	
50	88	
-88		
100.	1000.	0.

### 6.2.34 Control statement **node** (optional)

Specify the node numbers for which detailed output is desired.

Group 1 - M

Group 2 - MN (1), MN (2), . . . , MN (M)

Group 3 - X, Y, Z (as needed)

or

Group 1 - KEYWORD

Group 2 - JA, JB, JC (JA, JB, JC - defined on page 19)

Input variable	Format	Description
M	integer	Number of nodes for which information will be printed on the output (iout) and history plot (ishis, istr) files. If $M \leq 0$ , pressure and temperature will be written on the output file for all nodes, but no nodal parameter values will be printed in the history plot files. Group 2 is omitted if $M \leq 0$ .
MN	integer	M node numbers for which information will be printed on the output file (iout). If $MN(l) < 0$ , then coordinates are used to define the print-out node, and the coordinate sets (X, Y, Z) for each $MN(l) < 0$ are added after Group 2.
X	real	Coordinates of node for which information will be printed. One line for each $MN < 0$ . The code finds the node closest to the coordinate given. For 2-D problems, set $Z = 0$ . No input if no $MN > 0$ .
Y	real	
Z	real	
KEYWORD	character*5	Key word for invoking node specification by ja, jb, jc format. The necessary word is <i>block</i> .

The following are examples of **node**:

node
2
50      88

node
2
50      -88
100.    1000.    0.

node
block
1      100      10
-3      0      0

### 6.2.35 Control statement perm (required)

Assign permeabilities of the rock. Permeabilities represent average values of a volume associated with a node. Note that using **rlp** to describe relative permeabilities causes these values to be overwritten.

Group 1 - JA, JB, JC, PNXD, PNYD, PNZD (JA, JB, JC - defined on page 19)

Input variable	Format	Default	Description
PNXD	real	1.e-30	Permeability in the x-direction ( $m^2$ ).
PNYD	real	1.e-30	Permeability in the y-direction ( $m^2$ ).
PNZD	real	1.e-30	Permeability in the z-direction ( $m^2$ ).



The following is an example of **perm**:

perm					
1	140	1	2.50e-14	2.50e-14	0.00e-00

### 6.2.36 Control Statement pest (optional)

Out variable information for PEST parameter estimation routine.

Group 1 - MPEST

Group 2 - (NPEST(I), I=1, MPEST)

Group 3 - X, Y, Z (as needed)

Input Variable	Format	Description
MPEST	integer	Number of nodes for PEST output. At present the code outputs only pressures, saturations, and temperatures.
NPEST(I)	integer	Node numbers printed to the output file (suffix .pest) with values of variables listed above. If NPEST(I) < 0, then the node numbers are calculated with the coordinates.
X, Y, Z	real	Coordinates in grid if NODE(I) < 0. The coordinates are used to find the node closest in distance to that point, and that node is substituted for NODE(I).

### 6.2.37 Control statement ppwr (optional)

Group 1 - IPOROS

Group 2 - JA, JB, JC, POR1, POR2, POR3, POR4 (number depends on model type) (JA, JB, JC - defined on page 19)

Input variable	Format	Description
IPOROS	integer	Model type.  <b>Model (-1):</b> IPOROS = -1, input of specific storage $S_s$ .
POR1	real	Specific storage ( $m^{-1}$ ) $S_s = \rho g(\alpha + \phi\beta)(m^{-1})$ where $\rho$ = liquid density $g$ = gravity $\alpha$ = aquifer compressibility $\phi$ = porosity $\beta$ = liquid compressibility

Input variable	Format	Description
<b>Model (1):</b> IPOROS = 1, input of aquifer compressibility		
POR1	real	<p>Aquifer compressibility model.</p> $\phi = \phi_o + \alpha(P - P_o)$ <p>where</p> <p><math>\alpha</math> = aquifer compressibility</p> <p><math>\phi</math> = initial porosity</p> <p><math>P_o</math> = initial pressure</p>
<b>Model (-2):</b> IPOROS = -2, Gangi model with calculation of initial permeability and porosity.		
POR1	real	<p>Exponent <math>m</math> in Gangi bed-of-nails model.</p> $\phi = \phi_o \left[ 1 - \left( \frac{P_c}{P_x} \right)^m \right]$ $P_c = \sigma - P - \alpha E (T - T_o)$ <p>where</p> <p><math>\phi</math> = porosity</p> <p><math>\phi_o</math> = initial porosity</p> <p><math>m</math> = Gangi exponent</p> <p><math>P_x</math> = fitted parameter (MPa)</p> <p><math>\sigma</math> = in-situ stress (MPa)</p> <p><math>\alpha</math> = coefficient of thermal expansion <math>\left( \frac{1}{^\circ\text{C}} \right)</math></p> <p><math>E</math> = Young's modules (MPa)</p> <p><math>T</math> = temperatures (<math>^\circ\text{C}</math>)</p> <p><math>T_o</math> = initial temperature (<math>^\circ\text{C}</math>)</p>
POR2	real	$P_x$ parameters (MPa) in Gangi equation.
POR3	real	$\sigma$ , in-situ stress (Mpa).
POR4	real	<p><math>(\alpha E)</math> : the product of the coefficient of thermal expansion for the rock and the Young's modules (MPa/<math>^\circ\text{C}</math>). Note: for the Gangi model, the permeability is varied by <math>K = K_o \left( \frac{\phi}{\phi_o} \right)^3</math>.</p>

### 6.2.38 Control statement pres (required if macro init not used)

Group 1 - JA, JB, JC, PHRD, TIND, IEOSD (JA, JB, JC - defined on page 19)

The initial values defined in control statement **pres** supersede all others. Note that the term “saturated” is a thermodynamic definition and not the groundwater hydrology definition (volumetric fraction of pore void that is filled with water: IEOSD = 1). Saturated here indicates that vapor and liquid phases exist simultaneously. The superheated region means that all pore space is filled with gas.

Input variable	Format	Default	Description
PHRD	real	PEIN	Initial pressure (MPa).
TIND	real		Initial temperature (°C) if IEOSD = 1 or 3; initial saturation if IEOSD = 2
IEOSD	integer	1	Thermodynamic-region parameter. If IEOSD < 0, then the code uses ABS (IEOSD) and fixes the values of PHRD and TIND to the values provided above. IEOSD = 1, the compressed-liquid region. IEOSD = 2, the saturation region. IEOSD = 3, the superheated region.

The following is an example of **pres**:

pres					
-1	0	1	0.1	0.1	2
-2	0	1	0.1	0.1	2
-3	0	1	0.1	0.003	2
-4	0	1	0.1	0.1	2
-5	0	1	0.1	0.11	2
-6	0	1	0.1	0.11	-2
1	800	1	0.1	0.5	2

### 6.2.39 Control statement **ptrk** (optional, cannot be used with **trac**)

Group 1 - NPART, RSEED

Group 2 - DAYCS, DAYCF, DAYHF, DAYHS

Group 3 - TRAK\_TYPE, HALF\_LIFE, POUT, PRNT\_RST

Group 4 - TRANSFLAG(JJ), KD(JJ), TCLX(JJ), TCLY(JJ), TCLZ(JJ),  
DIFFMAT(JJ), RD\_FRAC(JJ), MATRIX\_POR(JJ), FSPACING(JJ)

Group 5 - JA, JB, JC, ITRC (JA, JB, JC - defined on page 19)

Group 6- JA, JB, JC, PCNSK, T1SK, T2SK (JA, JB, JC - defined on page 19)

Group 4 is used to define models in which identical sorption and transport parameters are assumed to apply. Group-4 data are read until a blank line is encountered. The model number JJ is incremented by 1 each time a line is read.

The concentration output is written to the .trc, .out, AVS concentration output files, and the .fin file, if specified (nonzero value of PRNT\_RST).

Input variable	Format	Description
NPART	integer	Number of particles in the simulation. Note: the actual number may be slightly less than the number specified by the user because when the code divides the particles among the starting nodes as specified in Group 7, the code must input an integer number of particles at each node.
RSEED	integer	6-digit integer random number seed.
DAYCS	real	Time that the particle-tracking solution is enabled (days).
DAYCF	real	Time that the particle-tracking solution is disabled (days).
DAYHF	real	Time that the flow solution is disabled (days).
DAYHS	real	Time that the flow solution is enabled (days).
TRAK_TYPE	integer	Flag to denote the fluid phase of the particles. 1 - liquid-phase particles. 2 - vapor-phase particles.
HALF_LIFE	real	Half-life for irreversible first-order decay reaction(s). Set HALF_LIFE = 0 for no decay.
POUT	integer	<p>Flag to specify the concentration output.</p> <p>1 - Concentrations computed as number of particles per unit total volume (rock and fluid).</p> <p>2 - Concentrations computed as number of particles per unit fluid volume (the fluid is liquid for TRAK_TYPE = 1 and gas for TRAK_TYPE = 2).</p> <p>3 - Concentrations computed as number of particles at a given node point.</p> <p>4 - Used for radioactive-particle mixing model (only liquid tracer). For meaningful results, the particles must all be injected simultaneously in a pulse (give a very short duration of injection starting at time 0). The file getconc.f contains data describing the function <math>f(t)</math> vs. time, where <math>f(t)</math> is given as</p> $\int_0^t C(t) \exp(-kt) dt$ <p>The meaningful output are the final concentrations after all the particles have left the system.</p> <p>-1, -2, -3, or -4 - Concentrations computed as specified above for abs(pout). The .trc file contains breakthrough output for the first node specified in the node macro.</p> <p>0 - Concentration output is a running total of the number of particles that have left each node divided by the fluid or vapor mass at that node, depending on trak_type.</p>

Input variable	Format	Description
PRNT_RST	integer	<p>Flag to specify whether particle information is written to the ".fin" file.</p> <p>0 - Particle information is not written to ".fin" file.</p> <p>1 - Particle information is written to the ".fin" file.</p> <p>-1 - Particle positions and ages are written to the ".fin" file.</p> <p>When particle-tracking data are written to the .fin file, the arrays are written after all of the heat- and mass-simulation information. The information written is sufficient to perform a restart of the particle-tracking simulation and to postprocess the data to compile statistics on the particle-tracking run. However, for a large number of particles, this file can become quite large, so particle-tracking information should only be written when necessary. Thus, 0 should be used for PRNT_RST unless restarting or postprocessing to obtain particle statistics is required. Selecting the -1 option allows a subset of the full set of information needed for a restart (particle positions and ages) to be written. Restart runs that use this file as input will only be approximate because the particle is assumed to have just entered its current cell. For restart runs, PRNT_RST = 1 is preferred, whereas PRNT_RST = -1 is appropriate for output of particle statistics for postprocessing.</p>
TRANSFLAG	integer	<p>Flag to specify which transport mechanisms apply.</p> <p>1 - advection only (no dispersion or matrix diffusion).</p> <p>2 - advection and dispersion (no matrix diffusion).</p> <p>3 - advection and matrix diffusion (no dispersion).</p> <p>4 - advection, dispersion, and matrix diffusion.</p>
KD	real	<p>Sorption coefficient (linear, reversible, equilibrium sorption). Units are kg-fluid/kg-rock (these units are equivalent to the conventional units of cc/g when the carrier fluid is water at standard conditions). This value applies to the medium as a whole when matrix diffusion is turned off, whereas for simulations invoking matrix diffusion, the value applies to the rock matrix. For the latter case, sorption in the flowing system (fractures) is modeled using the RD_FRAC variable.</p>
TCLX	real	<p>Dispersivity in the x-direction (m). The input value is ignored when dispersion is turned off.</p>
TCLY	real	<p>Dispersivity in the y-direction (m). The input value is ignored when dispersion is turned off.</p>
TCLZ	real	<p>Dispersivity in the z-direction (m). The input value is ignored when dispersion is turned off.</p>
DIFFMAT	real	<p>Molecular diffusion coefficient in the rock matrix (m<sup>2</sup>/s). The input value is ignored unless matrix diffusion is invoked.</p>
RD_FRAC	real	<p>Retardation factor within the primary porosity (fractures) for a matrix-diffusion particle-tracking simulation (use 1 for no sorption on fracture faces). The input value is ignored unless matrix diffusion is invoked.</p>

Input variable	Format	Description
MATRIX_POR	real	Porosity of the rock matrix. Used to simulate diffusion and sorption in the rock matrix when matrix diffusion is invoked. Note: when matrix diffusion is turned off, particle transport through the medium is computed using the porosity set in the <b>rock</b> macro, and the input value of MATRIX_POR is ignored.
FSPACING	real	Mean fracture spacing (m). When matrix diffusion is invoked, the mean fracture aperture (a parameter in the matrix-diffusion model) is computed as fracture porosity (from the <b>rock</b> macro) divided by FSPACING. When matrix diffusion is turned off, the value of FSPACING is ignored.
ITRC	integer	Model number for parameters defined in group 4. Default is [1].
PCNSK	real	<p>Particle-injection parameter assigned for nodes defined by JA, JB, and JC. Two options are available.</p> <p>PCNSK &gt; 0 - particles are injected at each node in proportion to the source mass flow rate at the node. When multiple lines of input are given for Group 6, PCNSK is proportional to the particle-injection concentration. This boundary condition is equivalent to injecting a solute of a given concentration into the system. Note: the source flow rates used to assign the number and timing of particle injections are those at the beginning of the particle-tracking simulation (time DAYCS). Transient changes in this source flow rate during the particle-tracking simulation do not change the input of particles to the system.</p> <p>PCNSK &lt; 0 - particles are introduced at the node(s), regardless of whether there is a fluid source at the node. When multiple lines of input are given for Group 6, abs(PCNSK) is proportional to the number of particles introduced at the node(s).</p> <p>When multiple lines of input are given for Group 6, all PCNSK values must have the same sign (i.e., the two options cannot be invoked in the same simulation). Default is 0 for all unassigned nodes, meaning that no particles are injected at that node.</p>
T1SK	real	Time (days) when particle injection begins. Default is [0].
T2SK	real	Time (days) when particle injection ends. Default is [0].

*Notes on Restarting:* As with all restart runs for FEHM, a .ini file is specified to be read to set the initial conditions upon restarting. However, there are two possibilities for restart calculations with particle tracking: 1) the heat- and mass-transfer solution is being restarted, but the particle-tracking simulation is initiated during the restart run (it was not carried out in the simulation that generated the .ini file); or 2) the heat- and mass-transfer solution and the particle-tracking simulation are both being restarted. If the code does not find the "ptrk" key word at the top of the .ini file, then the original run did not employ particle tracking, and Case 1 is assumed. A common example is a preliminary calculation that establishes a fluid-flow steady state, followed by a restart simulation of transport.

If "ptrk" was written into the .ini file in the original run, the particle data in the .ini file are read and used to initialize the particle-tracking simulation (Case 2). In this instance, the number of particles (NPART) must be set the same for the restart run as in the original run, or the results will be unpredictable. When restarting a particle-tracking simulation, certain input data are overwritten by information in the .ini file. These parameters include RSEED, PCNSK, T1SK, and T2SK. Other input parameters can be set to different values in the restart run than they were in the original run, but of course, care must be taken to avoid physically unrealistic assumptions, such as an abrupt change in transport properties of Group 4 part way through a simulation.

A final note on restart calculations is in order. A common technique in FEHM restart calculations is to reset the time at the top of the .ini file to 0, so that the starting time of the restart simulation is arbitrarily 0 rather than the ending time of the original simulation. This technique is useful for the example of the steady-state-flow calculation, followed by a restart solute-transport calculation. Although the technique is acceptable for particle-tracking runs that are initiated only upon restart (Case 1), it is invalid when a particle-tracking run is being resumed (Case 2). The reason is that all particle times read from the .ini file are based on the starting time of the original simulation during which the particle-tracking simulation was initiated.

The following is an example of ptrk:

ptrk								
100000	122945							
10.	20.	10.	20.					
1	0	2	0					
4	0.	2.	2.	2.	5.e-11	1.	0.1	0.333
4	3.	2.	2.	2.	1.e-10	1.	0.28	2.
1	0	0	1					
-2	0	0	2					
-3	0	0	1.	10.	10.0001			

In this example, 100,000 nondecaying, liquid-borne particles are introduced as a sharp pulse (from time 10 to 10.0001 days) with the injection fluid in zone 3 (an injection well defined in the **zone** macro preceeding **ptrk**). The particle-tracking simulation starts as the heat- and mass-transfer simulation is turned off at day 10, after having established a fluid-flow steady state. Two models are defined for assigning transport properties of the particles. All nodes are assigned to model 1, after which model-2 properties are assigned for zone 2. A combined advection, dispersion, and matrix-diffusion model is used for all nodes. However, sorption in the matrix occurs only for model 2 (which is zone 2 in this

simulation), and the matrix-transport properties (porosity, fracture spacing, and diffusion coefficient) differ for this model as well.

#### 6.2.40 Control statement renm (optional)

Renumbers the node numbers.

Group 1 - JA, JB, JC, IGD (JA, JB, JC - defined on page 19)

Input variable	Format	Description
IGD	integer	New node number for given node.

#### 6.2.41 Control statement rflx (optional)

Radiation-heat source term. Not implemented in this version. A negative heat flux indicates heat flow into the reservoir.

Group 1 - EMISS

Group 2- JA, JB, JC, QFLUX, QFLXM (JA, JB, JC - defined on page 19)

Input Variable	Format	Description
EMISS	real	Emissivity.
QFLUX	real	If QFLXM = 0, then QFLUX is the heat flux (MW). If QFLXM ≠ 0, then QFLUX is a temperature, and the heat flux is calculated according to the formula: $Q_H = QFLXM(TL - QFLUX)$ (MW). [0]
QFLXM	real	Multiplier for heat-flux equation given in QFLUX description (MW/°C). If QFLXM = 0, then QFLUX is the heat flux (MW). [0]

#### 6.2.42 Control statement rlp (optional)

Relative-permeability and capillary-pressure model. Four models are available. The fifth model flag is used to designate that **rlp** data should be read from an auxiliary file.

Group 1 - IRLP(i), RP1, RP2, RP3, RP4, RP5, RP6, RP7, RP8, RP9, RP10, RP11, RP12, RP13, RP14, RP15 (number of parameters depends on model selected)

Group 2 - JA, JB, JC, I (JA, JB, JC - defined on page 19)

Only those parameters defined for a given model need to be input. Group 1 is ended when a blank line is encountered. The parameter (i) is incremented each time a Group 1 line is read. Group 2 lines will refer to this parameter. For model number 4 (the combined van Genuchten model), the permeability is isotropic and overwrites the input from macro **perm**.

Or: Group 1 -IRLP(i)

Group 2 - RLPFILE



Input variable	Format	Description
IRLP(i)	integer	Relative-permeability model type.
<b>Model 1:</b> IRLP(i) = 1, linear relative permeability, linear capillary pressure (6 parameters required).		
RP1	real	Residual liquid saturation.
RP2	real	Residual vapor saturation.
RP3	real	Maximum liquid saturation.
RP4	real	Maximum vapor saturation.
RP5	real	Capillary pressure at zero saturation (Mpa).
RP6	real	Saturation at which capillary pressure goes to zero.
<b>Model 2:</b> IRLP(i) = 2, Corey relative permeability, linear capillary pressure (4 parameters required).		
RP1	real	Residual liquid saturation.
RP2	real	Residual vapor saturation.
RP3	real	Capillary pressure at zero saturation (MPa).
RP4	real	Saturation at which capillary pressure goes to zero.
<b>Model 3:</b> IRLP(i) = 3, van Genuchten relative permeability, van Genuchten capillary pressure (6 parameters required).		
RP1	real	Residual liquid saturation.
RP2	real	Maximum liquid saturation.
RP3	real	Inverse of air entry pressure, $\alpha_G$ (1/m). (Note: some data are given in (1/Pa); convert using pressure = $\rho g \Delta h$ .)
RP4	real	Power n in van Genuchten formula.
RP5	real	Low-saturation fitting parameter, multiple of cutoff capillary pressure assigned as maximum capillary pressure. If $RP5 < 0$ , then a linear fit from this cutoff saturation (RP6 is used). The slope of the cutoff saturation is used to extend the function to saturation = 0. If $RP5 = 0$ , a cubic fit is used. The slope at the cutoff saturation is matched, and the conditions $\frac{\partial}{\partial S} P_{cap} = 0$ and $\frac{\partial^2}{\partial S^2} P_{cap} = 0$ are forced at $S = 0$ . If $RP5 > 0$ , a multiple of the value of the capillary pressure at the cutoff saturation, $RP5 \cdot P_{cap}(S_{cutoff})$ , is forced at $S = 0$ .
RP6	real	Cutoff saturation used in fits described for RP5. Must be greater than RP1.

Input variable	Format	Description
<b>Model 4:</b> IRLP(i) = 4, van Genuchten relative permeability, van Genuchten capillary pressure, effective continuum (15 parameters required).		
RP1	real	Residual liquid saturation, matrix rock material.
RP2	real	Maximum liquid saturation, matrix rock material.
RP3	real	Inverse of air entry pressure, $\alpha_G$ (1/m), matrix rock material. (Note: some data are given in (1/Pa); convert using pressure = $\rho g \Delta h$ .)
RP4	real	Power n in van Genuchten formula, matrix rock material.
RP5	real	Low-saturation fitting parameter, matrix rock material, multiple of cutoff capillary pressure assigned as maximum capillary pressure. If RP5 < 0, then a linear fit from this cutoff saturation (RP6 is used). The slope of the cutoff saturation is used to extend the function to saturation = 0. If RP5 = 0, a cubic fit is used. The slope at the cutoff saturation is matched, and the conditions $\frac{\partial}{\partial S} P_{cap} = 0$ and $\frac{\partial^2}{\partial S^2} P_{cap} = 0$ are forced at $S = 0$ . If RP5 > 0, a multiple of the value of the capillary pressure at the cutoff saturation, $RP5 \cdot P_{cap}(S_{cutoff})$ is forced at $S = 0$ .
RP6	real	Cutoff saturation used in fits described for RP5, matrix rock material. Must be greater than RP1.
RP7	real	Residual liquid saturation, fracture material.
RP8	real	Maximum liquid saturation, fracture material.
RP9	real	Inverse of air entry pressure, $\alpha_G$ (1/m), fracture material. (Note: some data are given in (1/Pa); convert using pressure = $\rho g \Delta h$ .)
RP10	real	Power n in van Genuchten formula, fracture material.
RP11	real	Low-saturation fitting parameter, fracture material, multiple of cutoff capillary pressure assigned as maximum capillary pressure. If RP11 < 0, then a linear fit from this cutoff saturation (RP6 is used). The slope of the cutoff saturation is used to extend the function to saturation = 0. If RP11 = 0, a cubic fit is used. The slope at the cutoff saturation is matched, and the conditions $\frac{\partial}{\partial S} P_{cap} = 0$ and $\frac{\partial^2}{\partial S^2} P_{cap} = 0$ are forced at $S = 0$ . If RP11 > 0, a multiple of the value of the capillary pressure at the cutoff saturation, $RP11 \cdot P_{cap}(S_{cutoff})$ , is forced at $S = 0$ .
RP12	real	Cutoff saturation used in fits described for RP11, fracture material. Must be greater than RP7.

Input variable	Format	Description
RP13	real	Fracture permeability ( $\text{m}^2$ ). This is the permeability of the individual fractures. The bulk permeability of the fracture continua is $RP13 \times RP15$ .
RP14	real	Matrix-rock saturated permeability ( $\text{m}^2$ ).
RP15	real	Fracture porosity.
<b>Model 5:</b> IRLP(i) = 5, a file containing the relative-permeability and capillary models is required. The name of this file is given on the line following IRLP(i) = 5. A model must be entered for each and every node in numerical (nodal) order. The models above are available. (See example.)		
I	integer	Number referring to the sequence of models read in Group 1. The default is [1].
RLPFILE	character*100	Name of the optional <b>rlp</b> data file.

The following are examples of **rlp**:

rlp				
2	0.3	0.1	2.0	1.
1	140	1	1	

rlp	
5	
rlp.dat	

File rlp.dat:

2	0.3	0.1	2.	1.
2	0.3	0.1	2.	1.
.	.	.	.	.
.	.	.	.	.
.	.	.	.	.
2	0.3	0.1	2.	1.

A model is  
entered for  
each node

### 6.2.43 Control statement rock (required)

Assign rock density, specific heat, and porosity.

Group 1 - JA, JB, JC, DENRD, CPRD, PSD (JA, JB, JC - defined on page 19)

Input variable	Format	Description
DENRD	real	Rock density (kg/m <sup>3</sup> ).
CPRD	real	Rock specific heat ( $\frac{\text{MJ}}{\text{kg} \cdot \text{K}}$ ). If CPRD > 1, the code will assume the units are ( $\frac{\text{J}}{\text{kg} \cdot \text{K}}$ ) and multiply by 10 <sup>-6</sup> .
PSD	real	Porosity.

The following is an example of **rock**:

rock					
1	140	1	2563.	1010.	0.3500

#### 6.2.44 Control statement rxn (optional)

Chemical reactions between species are invoked with this control statement. It is used in conjunction with control statement **trac**.

Group 1 - KEY\_GROUP

Group 2 - NGROUPS

Group 3 - GROUP(ISPECIES), ISPECIES = 1, NSPECI (repeated NGROUPS times, once for each group)

Group 4 - NRXNS, RXN\_INTERVAL

For each reaction, there are two possibilities that require different input: equilibrium reaction (KEY\_RXN = "equilibrium") and kinetic reaction (KEY\_RXN = "kinetic"). These two choices may be mixed in a given simulation to simulate a combination of equilibrium and kinetic reactions. Group-5 parameters are repeated NRXN times, once for each reaction. Within an equilibrium-reaction option, there are also two options based on the choice of the EQUIL\_MODEL parameter. For EQUIL\_MODEL = 1, the following input is used:

Group 5 - KEY\_RXN, EQUIL\_MODEL, EQUIL\_CONST25, ENTHALPY25, GAMMA\_CHECK, RATE\_FACTOR, ROUND\_TOL

For EQUIL\_MODEL = 2, the input is:

Group 5 - KEY\_RXN, EQUIL\_MODEL, AWWA(1), AWWA(2), AWWA(3), AWWA(4), AWWA(5), GAMMA\_CHECK, RATE\_FACTOR, ROUND\_TOL

Finally, for a kinetic reaction, the input is:

Group 5- KEY\_RXN, AR\_FOR, EA\_FOR, AR\_REV, EA\_REV

Group 6 - STOIC(ISPECIES), ISPECIES = 1, NSPECI (repeated NRXN times, once for each reaction)

- Group 7 - RATE\_POWER(ISPECIES), ISPECIES = 1, NSPECI (repeated NRXN times, once for each reaction)
- Group 8 - FL\_MULT(ISPECIES), ISPECIES = 1, NSPECI (repeated NRXN times, once for each reaction)
- Group 9 - SB\_MULT(ISPECIES), ISPECIES = 1, NSPECI (repeated NRXN times, once for each reaction)
- Group 10 - H\_MULT(ISPECIES), ISPECIES = 1, NSPECI (repeated NRXN times, once for each reaction)

Input for Groups 8 and 9 may be omitted by entering a blank line after the entries for Group 7. In this case, all reactions are assumed to occur with both fluid- and sorbed-phase solutes. If this is the case, or if there are no sorbing solutes in the simulation, then Groups 8 and 9 need not be entered. To specify the nature of the reactions involving sorbed-phase solutes (i. e., whether to compute the rate based on fluid concentration, sorbed-phase concentration, or both), see the variable descriptions below. Note that if any values in Group 8 or 9 are to be set other than to their default values, then all values for these two groups are required.

If there are no Henry's Law species present in the simulation, then Group 10 should be omitted. However, this group is necessary when a Henry's Law species is present, even if it does not participate in any reactions.

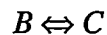
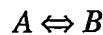
Input variable	Format	Description
KEY_GROUP	character	Key word to specify that species are to be placed into groups that are solved simultaneously. If "gr" is found as the first two characters of the line immediately after the rxn control statement, then the code reads in the species group information (Groups 2 and 3) of the input. Otherwise, the code assumes that each species is to be solved separately with an outer iterative loop to ensure overall convergence.
NGROUPS	integer	Number of groups of species. The species assigned to a group (using the GROUP input below) are solved simultaneously. If there is only one group, the solution is complete when convergence is achieved. If there is more than one group, an outer iterative loop over all groups is employed to ensure overall convergence.
GROUP	integer	NSPECIES values are entered for each line of input, and NGROUPS lines of input are required, one for each group. If a value is nonzero, then that species is present in the group. A value of zero denotes that the species is not present in the group. No more than four species can be assigned to any group, but there is no restriction on the number of groups that a species can be assigned to. Grouping of species that take part in rapid kinetics or equilibrium reactions is required for convergence. However, memory requirements increase as the square of the maximum number of species in a group.
NRXNS	integer	Number of chemical reactions.

Input variable	Format	Description
RXN_INTERVAL	real	This parameter allows the user to choose between two different iteration schemes as follows(see "Models and Methods Summary" (Zyvoloski et al. 1997) for a description of these solution procedures). RXN_INTERVAL = 0 - Solution scheme 1 always used. RXN_INTERVAL > 0 - Solution scheme 2 will be used RXN_INTERVAL times between each use of scheme 1.
KEY_RXN	character	Denotes the type of reaction. The first letter of the keyword is all that is required, but it must appear as the first character of the line. "equilibrium" - equilibrium reaction. "kinetic" - kinetic reaction.
EQUIL_MODEL	integer	Flag denoting which model is to be used for defining the temperature dependence of the equilibrium constant. 1 - van't Hoff model. 2 - multiparameter fit to experimental data for carbonate system.
EQUIL_CONST25	real	The term $A_{rxn}$ in equilibrium constant-temperature-dependence model.
ENTHALPY25	real	The term $\Delta H_H$ in equilibrium constant-temperature-dependence model.
GAMMA_CHECK	real	Equilibrium tolerance parameter $\gamma_{tol}$ . The reaction will be required to be at equilibrium to within $\gamma_{tol}$ at every node. For example, a value of $10^{-2}$ (recommended value) means that the reaction is at least 99% to equilibrium everywhere.
RATE_FACTOR	real	Parameter for scaling the rate constants used in the code for simulating equilibrium behavior. Recommended value: $10^{-3}$ .
ROUND_TOL	real	Cut-off parameter for forward reaction rate below which the check for equilibrium behavior is not made. Recommended value: $10^{-10}$ .
AWWA(1)	real	The term $A_{rxn,1}$ in equilibrium constant-temperature-dependence model.
AWWA(2)	real	The term $A_{rxn,2}$ in equilibrium constant-temperature-dependence model.
AWWA(3)	real	The term $A_{rxn,3}$ in equilibrium constant-temperature-dependence model.
AWWA(4)	real	The term $A_{rxn,4}$ in equilibrium constant-temperature-dependence model.
AWWA(5)	real	The term $A_{rxn,5}$ in equilibrium constant-temperature-dependence model.

Input variable	Format	Description
AR_FOR	real	Pre-exponential factor of the forward reaction (Eqn. 85 in Zyvoloski et al. (1997)). In keeping with the conventional method of defining rate constants, this parameter has units of $[(\text{concentration units})^p \times \text{s}]^{-1}$ , where $p$ is the sum of the exponents on all concentrations in the forward reaction minus 1. Thus, the units of the reaction rate are (concentration units)/s.
EA_FOR	real	Activation energy of the forward reaction (J/mol).
AR_REV	real	Pre-exponential factor of the reverse reaction.
EA_REV	real	Activation energy of the reverse reaction.
STOIC	real	For each solute, the stoichiometric coefficient (the $a$ 's in Eqn. 83 in Zyvoloski et al. (1987)) for the particular reaction. If positive, the solute is a reactant; if negative, the solute is a product; and if 0, the solute is not present in the reaction.
RATE_POWER	real	For each solute, the exponent in the rate law (the $b$ 's in Eqn. 84 in Zyvoloski et al. (1987)) for the particular reaction. If the corresponding value of STOIC is positive (the solute is a reactant), then $b = \text{RATE\_POWER}$ ; if the corresponding value of STOIC is negative (the solute is a product), then $b = -\text{RATE\_POWER}$ ; and if the corresponding value of STOIC is 0, this value of RATE_POWER is ignored. The reason for this convention is that when the law of microscopic reversibility applies (the exponents in the rate law are the stoichiometric coefficients), the values of STOIC and RATE_POWER are identical. Note also that a value of 0 for RATE_POWER when the corresponding value of STOIC is nonzero implies a zero-th order reaction and does not mean that the reactant is absent from the reaction.
FL_MULT	real	For each solute, a parameter signifying whether this particular reaction occurs for this solute in the fluid phase. Set the parameter to a nonzero value if the solute in the fluid phase is involved in the reaction, otherwise, set the parameter to 0. If this solute is not present in the reaction (the corresponding value of STOIC is 0), then the value of FL_MULT is irrelevant.

Input variable	Format	Description
SB_MULT	real	For each solute, a parameter signifying whether this particular reaction occurs for this solute in the sorbed phase. Set the parameter to a nonzero value if the solute in the sorbed phase is involved in the reaction, otherwise set the parameter to 0. If this solute is not present in the reaction (the corresponding value of STOIC is 0), then the value of SB_MULT is irrelevant.
H_MULT	real	For each Henry's Law species, set this parameter to 1 if the liquid-borne fraction of the solute is reacting or to -1 if the vapor-borne portion of the species is reacting. For each non-Henry's Law species, this parameter is not used but is read to keep a similar input format to SB_MULT and FL_MULT. Therefore, H_MULT <i>must</i> be set for <i>each</i> species in <i>every</i> reaction if a Henry's Law species is present in the simulation. Both the liquid- and vapor-borne portions of a Henry's Law species cannot react in one reaction. However, <i>two</i> reactions, one with the liquid portion reacting and the other with the vapor portion reacting, allow for both phases of the Henry's Law species to react.

The following example, along with the example data input for the **trac** macro, defines the reactive transport problem that is called Run 1 in Section 9.5. The reaction system contains three species (A, B, and C in the description below refer to species 1, 2, and 3, respectively) and the following two chemical reactions.



The reactions are reversible, with stoichiometric coefficients of 1, and the powers in the kinetic or equilibrium expressions are all 1 as well. Equilibrium sorption is specified for solute 1 (A) using the isotherm formulations in the **trac** macro. The FL\_MULT and SB\_MULT input are set so that the chemical reactions here pertain only to solute present in the fluid phase. Group 10 is omitted because for this example there are no Henry's Law species.

The first reaction is kinetically controlled, and forward and reverse rate constants (with no temperature dependence) are given. The second reaction is an equilibrium reaction, and an equilibrium constant (with no temperature dependence) is given. The code will iteratively solve the system in two groups, first with A only, then with B and C coupled. Coupling of the solutes in the equilibrium reaction is necessary for convergence, but solutes that are uncoupled or weakly coupled through "slow" kinetic reactions can be solved separately to minimize computer storage requirements. If the number of finite-element nodes is small enough, all three species could be coupled, and better performance would be expected if a large number of outer iterations are required. To do this in the example above, NGROUP would be set to 1, and a single line of three



1's for GROUP would denote that all three solutes are to be solved simultaneously.

The following is the example of **rxn**:

rxn						
group						
2						
1	0	0				
0	1	1				
2	0					
kinetic	3.1688e-11	0.	3.1688e-10	0.		
equilibrium	1	0.2	0.	1.e-2	1.e-3	1.e-10
1	-1	0				
0	1	-1				
1	-1	0				
0	1	-1				
1	1	1				
1	1	1				
0	0	0				
0	0	0				

## 6.2.45 Control statement **sol** (required)

Group 1 - NTT, INTG

Input variable	Format	Description
NTT	integer	Parameter that defines the type of solution required. NTT > 0 - coupled solution. NTT ≤ 0 - heat-transfer-only solution.
INTG	integer	Parameter that defines element integration type. INTG ≤ 0 - Lobatto (node point) quadrature is used, recommended for heat and mass problems without stress. INTG > 0 - Gauss quadrature is used, recommended for problems requiring a stress solution. This is the default for 3-D problems.

The following is an example of **sol**:

sol	
1	-1

## 6.2.46 Control statement **solv** (Not implemented)

## 6.2.47 Control statement **stea** (optional)

No input is associated with this macro statement. The code will attempt to find a steady-state solution if present. Not tested.

This statement enables a 1-D solution in the y-direction (2-D) or z-direction (3-D) when gravity is present to generate an initial steady-state solution.

### 6.2.48 Control statement stop (required)

No input is associated with this control statement. It signals the end of input, and as such, it always appears as the last line of an input deck.

### 6.2.49 Control statement strs (Not implemented in this version of FEHM)

### 6.2.50 Control Statement svar (optional)

Group 1- IVAR

Input variable	Format	Description
IVAR	integer	If IVAR =1, then pressure-enthalpy variables used in place of pressure-temperature variables for water/water-vapor physics. Otherwise not enabled.

### 6.2.51 Control statement text (optional)

Group 1- WDD1

Input variable	Format	Description
WDD1	character*80	Line of text. A maximum of 80 characters per line are entered. Text is input until a blank line is inserted to signal the end of the control statement. This text is written to the output file (iout).

The following is an example of **text**:

```
text
This is a 2-D model of the PACE problem
It will be used to study thermal effects
user # = -20 to get waste packages
```

### 6.2.52 Control statement thic (optional)

Input for variable thickness for two-dimensional problems.

Group 1 - JA, JB, JC, THIC (JA, JB, JC - defined on page 19)

Input variable	Format	Description
THIC	real	Thickness of the model domain in the third dimension (m). Default is [1].

The following is an example of **thic**:

```
thic
1      0      0      10.
-2     0      0      5.
```

In this example, the thickness for all nodes is set to 10 m, after which the nodes defined by zone 2 are set to 5 m. Thus, the thickness is 10 m everywhere except zone 2, where thickness is 5 m.

### 6.2.53 Control statement time (required)

Time-step and time-of-simulation data.

Group 1 - DAY, TIMS, NSTEP, IPRTOUT, YEAR, MONTH, INITTIME

Group 2 - DIT1, DIT2, DIT3, ITC (as needed)

DAY should be larger than DAYMIN defined in Control Statement **ctrl**. The code proceeds to the next control statement when a blank line is encountered for Group 2. This can be used to generate output at specific times (with multiple group 2s). A contour plot will be drawn at each DIT1 regardless of the input in control statement **cont**. The restart file will be written (or rewritten if one already exists) at each DIT1.

Input variable	Format	Description
DAY	real	Initial time-step size (days).
TIMS	real	Final simulation time (days).
NSTEP	integer	Maximum number of time steps allowed.
IPRTOUT	integer	Print-out interval for nodal information (pressure, enthalpy etc.), as set up under control statement node (i.e., number of time steps).
YEAR	integer	Year that simulation starts.
MONTH	integer	Month that simulation starts.
INITTIME	real	Initial time of simulation (days). For compatibility with older versions, if this parameter is absent, the initial time of simulation will be 0 if no restart file is used or the time in the restart file if one is used.
DIT1	real	Time (days) for time-step change.
DIT2	real	New time-step size (days). If $DIT2 < 0$ , then $ABS(DIT2)$ is the new time-step multiplier.
DIT3	real	Implicitness factor for new time step. DIT3 $\leq$ 1.0 - backward Euler. DIT3 $>$ 1.0 - second-order implicit scheme.
ITC	integer	New print-out interval.

The following is an example of **time**:

time						
30.0	3650.0	20	5	1989	10	0.0
1.0	-1.2	1.0	10			

### 6.2.54 Control statement trac (optional)

Group 1 - USER\_MACRO, ANO, AWC, EPC, UPWGTA

Group 2 - DAYCS, DAYCF, DAYHF, DAYHS

Group 3 - IACCMX, DAYCM, DAYCMM, DAYCMX

Group 4 - NSPECIES

Group 5 - INPUT\_MSG

There are two options for group six. If the same diffusion coefficient and dispersivities are to be used for all species of the same type (liquid and/or vapor), then a keyword must be entered. This will make the calculations more efficient and thus should be used if applicable. In the absence of a keyword, the following input is used:

Group 6 - ICNS

Group 7 - IADSF, A1ADSF, A2ADSF, BETADF, DIFFM, TCX, TCY, TCZ

Group 8 - JA, JB, JC, ITRCD (JA, JB, JC - defined on page 19)

Group 9 - HENRY\_MODEL, HAWWA(1), HAWWA(2), HAWWA(3),  
HAWWA(4), HAWWA(5) (only for a Henry's Law species)

Group 10 - JA, JB, JC, ANQO (JA, JB, JC - defined on page 19)

Group 11 - JA, JB, JC, CNSK, T1SK, T2SK (JA, JB, JC - defined on page 19)

Groups 6, 7, 8, 9, 10, and 11 are entered as a unit for each solute. However, for a solid species, only groups 6, 10, and 11 are entered (groups 7, 8, and 9 are not applicable).

Groups 7 and 8 are used to define transport models for which sorption and dispersion parameters are identical. For a liquid or vapor species, only one set of Group 7 parameters should be entered per region. However, for a Henry's Law species, two sets of parameters per region must be entered. For this case, the liquid sorption parameters should be entered on the first line and the vapor sorption parameters on a second line or as a continuation of the first line. Groups 7 and 8 are not applicable to a solid species and should not be entered. Group 7 is read until a blank line is encountered. The model number is incremented by 1 each time a line is read. Group 8 then assigns a transport model number to every node.

For the same diffusion coefficient and dispersivities:

Group 6 - KEYWORD

If only liquid species are present (keyword *dspl*) or only vapor species are present (keyword *dspv*), Group 7 is defined as follows:

Group 7 - DIFFM, TCX, TCY, TCZ

Otherwise, if both liquid and vapor are present (keyword *dspb*), parameters for both must be entered:

Group 7- DIFFML, TCXL, TCYL, TCZL, DIFFMV, TCXV, TCYV, TCZV

Group 8 - JA, JB, JC, ITRCDSP (JA, JB, JC - defined on page 19)

Group 9 - ICNS

Group 10 -IADSF, A1ADSF, A2ADSF, BETADF

Group 11 -JA, JB, JC, ITRCD (JA, JB, JC - defined on page 19)

Group 12 -HENRY\_MODEL, HAWWA(1), HAWWA(2), HAWWA(3),  
HAWWA(4), HAWWA(5) (only for a Henry's Law species)

Group 13 -JA, JB, JC, ANQO (JA, JB, JC - defined on page 19)

Group 14 - JA, JB, JC, CNSK, T1SK, T2SK (JA, JB, JC - defined on page 19)

Injection nodes must be specified in control statement **flow**.

Input variable	Format	Description
USER_MACRO	character*5	Key word for invoking a solute transport user subroutine. If the word <i>userc</i> is placed in this position, then the code invokes a solute-transport user subroutine at each time step. Omit this key word if there is no solute user subroutine for the simulation.
ANO	real	Initial solute concentration set at all nodes for all species unless overwritten by a restart file input or values in group 9 below (moles/kg fluid).
AWC	real	Implicitness factor for solute solution. AWC > 1.0 gives 2nd-order solution. AWC ≤ 1.0 gives 1st-order solution.
EPC	real	Equation tolerance for solute solution.
UPWGTA	real	Upstream weighting term for the solute solution. UPWGTA < 0.5 - UPWGTA is set to 0.5. UPWGTA > 1.0 - UPWGTA is set to 1.0.
DAYCS	real	Time that the solute solution is enabled (days).
DAYCF	real	Time that the solute solution is disabled (days).
DAYHF	real	Time that the flow solution is disabled (days).
DAYHS	real	Time that the flow solution is enabled (days).
IACCMX	integer	Maximum number of iterations allowed in solute solution if time-step multiplier is enabled.
DAYCM	real	Time-step multiplier for solute solution.
DAYCMM	real	Initial time step for solute solution (days).
DAYCMX	real	Maximum time step for solute solution (days).
NSPECIES	integer	Number of solutes simulated.

Input variable	Format	Description
INPUT_MSG	character*4	Keyword ' <i>ldsp</i> ' specifying longitudinal/transverse dispersion. If x-, y-, z-dispersion is desired, Group 5 is omitted, and dispersivities are input in x, y, and then z order. Otherwise, if longitudinal/transverse dispersion is desired, the keyword ' <i>ldsp</i> ' is entered, and dispersivities are instead input in longitudinal and then transverse order with values for the third dimension omitted.
KEYWORD	character*4	Keyword specifying the same diffusion coefficient and dispersivities are to be used for all species of the same type (liquid and/or vapor). ' <i>dspI</i> ' indicates that only liquid species exist. ' <i>dspv</i> ' indicates that only vapor species exist. ' <i>dspb</i> ' indicates that both liquid and vapor species exist.
ICNS	integer	Phase designation for the <i>ith</i> solute. -2 - Henry's Law species (input and output concentration values are gas concentrations). -1 - Vapor species. 0 - Solid species. 1 - Liquid species. 2 - Henry's Law species (input and output concentration values are liquid concentrations).
IADSF	integer	Adsorption model type for the <i>ith</i> species, <i>ith</i> region. 0 - conservative solute. 1 - linear sorption isotherm. 2 - Freundlich sorption isotherm. 3 - Modified Freundlich sorption isotherm. 4 - Langmuir sorption isotherm.
A1ADSF	real	$\alpha_1$ parameter in adsorption model.
A2ADSF	real	$\alpha_2$ parameter in adsorption model.
BETADF	real	$\beta$ parameter in adsorption model.
DIFFM	real	Molecular diffusion coefficient ( $m^2/s$ ).
DIFFML	real	Molecular diffusion coefficient for liquid ( $m^2/s$ ).
DIFFMV	real	Molecular diffusion coefficient for vapor ( $m^2/s$ ).
TCX	real	Dispersivity in x-direction (m).
TCY	real	Dispersivity in y-direction (m).
TCZ	real	Dispersivity in z-direction (m).
TCXL	real	Dispersivity in x-direction for liquid (m).
TCYL	real	Dispersivity in y-direction for liquid (m).
TCZL	real	Dispersivity in z-direction for liquid (m).
TCXV	real	Dispersivity in x-direction for vapor (m).
TCYV	real	Dispersivity in y-direction for vapor (m).

Input variable	Format	Description
TCZV	real	Dispersivity in z-direction for vapor (m).
ITRCD	integer	Region number for sorption and dispersion parameters given in group 7 (no keyword) or for sorption parameters given in group 11. Default is [1].
ITRCDSP	integer	Region number for dispersion parameters given in group 7 (keyword). Default is [1].
HENRY_MODEL	integer	Flag denoting which model to be used for defining the temperature dependence of the Henry's Law constant. 1 - van't Hoff model. 2 - multiparameter fit to experimental data (used for carbonate system).
HAWWA(1)	real	Term in Henry's Law temperature-dependence model. For model 1 - parameter value is $A_H$ . For model 2 - parameter value is $A_{H,1}$ .
HAWWA(2)	real	Term in Henry's Law temperature-dependence model. For model 1 - parameter value is $\Delta H_H$ . For model 2 - parameter value is $A_{H,2}$ .
HAWWA(3)	real	Term in Henry's Law temperature-dependence model. For model 1 - not used. For model 2 - parameter value is $A_{H,3}$ .
HAWWA(4)	real	Term in Henry's Law temperature-dependence model. For model 1 - not used. For model 2 - parameter value is $A_{H,4}$ .
HAWWA(5)	real	Term in Henry's Law temperature-dependence model. For model 1 - not used. For model 2 - parameter value is $A_{H,5}$ .
ANQO	real	Initial concentration of tracer, which will supercede the value given by ANO in group 1. Note that if initial values are read from a restart file, these values will be overwritten. Units are moles per kg vapor or liquid for a liquid, vapor, or Henry's Law species and moles per kg of solid for a solid species. Default is [0].
CNSK	real	Injection concentration at inlet node (moles per kg liquid or vapor). If fluid is exiting at a node, then the in-place concentration is used. If $CNSK < 0$ , then the concentration at that particular node will be held at a concentration of $abs(CNSK)$ . Default is [0] for all unassigned nodes.
T1SK	real	Time (days) when tracer injection begins. Default is [0].
T2SK	real	Time (days) when tracer injection ends. Default is [0].

In the following example of **trac**, three liquid solutes are simulated. The solute transport solution is turned on as the heat and mass solution is turned off at day 3.6525e6. Solute 1 sorbs with an equilibrium sorption  $K_d$  value of 0.1; solutes 2 and 3 exhibit no sorption. All three solutes have the same transport parameters, although this is not a requirement of the code, even when the solutes are coupled through chemical reactions. All solutes start at a concentration of 0 within the model (in this case, a one-dimensional column). Solute 1 is injected at a concentration of 1 for a short time interval; there is no source for solutes 2 or 3. This example is meant to be used in combination with the example given for the **rxn** macro, which defines a system of chemical reactions among the three solutes. Therefore, solutes 2 and 3 are generated only through chemical reactions (neither of these solutes appear in the system initially, and there is no injection source term for either solute). The second example of **trac** is modified for use with longitudinal and transverse dispersion. The third and fourth examples are the above two examples of **trac**, modified to illustrate keyword use for the same diffusion coefficient and dispersivities. The reactive transport problem specified here and in the example **rxn** macro is discussed further in Section 9.5.

trac								
0	1	1.e-6	.5					
3.6525e6	5.47875e6	3.6525e6	5.47875e6					
50	1.2	3.6525e3	3.6525e4					
3								
1								
1	0.1	0.	1.	1.e-10	50	1.e-30	1.e-30	
1	0	0	1					
1	0	0	0.					
1	202	201	1.	3.6525e6	3.689025e6			
1								
0	0	0	1.	1.e-10	50	1.e-30	1.e-30	
1	0	0	1					
1	0	0	0.					
1								
0	0	0	1.	1.e-10	.033333	1.e-30	1.e-30	
1	0	0	1					
1	0	0	0.					



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trac						
0	1	1.e-6	.5			
3.6525e6	5.47875e6	3.6525e6	5.47875e6			
50	1.2	3.6525e3	3.6525e4			
3						
ldsp						
1						
1	0.1	0.	1.	1.e-10	50	5.0
1	0	0	1			
1	0	0	0.			
1	202	201	1.	3.6525e6	3.689025e6	
1						
0	0	0	1.	1.e-10	50	5.0
1	0	0	1			
1	0	0	0.			
1						
1						
0	0	0	1.	1.e-10	.0333	.00333
1	0	0	1			
1	0	0	0.			

trac						
0	1	1.e-6	.5			
3.6525e6	5.47875e6	3.6525e6	5.47875e6			
50	1.2	3.6525e3	3.6525e4			
3						
dspl						
1.e-10	.0333333	1.e-30	1.e-30			
-1	0	0	1			
1						
1	0.1	0.	1.			
1	0	0	1			
1	0	0	0.			
1	202	201	1.	3.6525e6	3.689025e6	
1						
0	0	0	1.			
1	0	0	1			
1	0	0	0.			

1			
0	0	0	1.
1	0	0	1
1	0	0	0.

trac					
0	1	1.e-6	.5		
3.6525e6	5.47875e6	3.6525e6	5.47875e6		
50	1.2	3.6525e3	3.6525e4		
3					
ldsp					
dspl					
1.e-10	.0333	.00333			
-1	0	0	1		
1					
1	0.1	0.	1.		
1	0	0	1		
1	0	0	0.		
1	202	201	1.	3.6525e6	3.689025e6
1					
0	0	0	1.		
1	0	0	1		
1	0	0	0.		
1					
0	0	0	1.		
1	0	0	1		
1	0	0	0.		

### 6.2.55 Control statement user (optional)

Group 1 - KK

Input variable	Format	Description
KK	integer	Integer number passed to subroutine <b>user</b> for user-defined input parameters. This user subroutine call differs from the one invoked in the control file in that whereas that subroutine is called at every time step, this one is called only at the beginning of the simulation to set parameters that do not change later in the simulation.

### 6.2.56 Control statement vcon (optional)

Group 1 - IVCON(i), VC1F(i), VC2F(i), VC3F(i)

Group 2 - JA, JB, JC, IVCND (JA, JB, JC - defined on page 19)

The parameter (i) is incremented each time Group 1 is read. Group 2 lines will refer to this parameter. Group 1 is ended with a blank line.

Input variable	Format	Description
IVCON(i)	integer	Model type for ith conductivity model. IVCON(i) = 1 - linear variation of thermal conductivity with temperature. IVCON(i) = 2 - square-root variation of thermal conductivity with liquid saturation.
VC1F(i)	real	Reference temperature ( $^{\circ}\text{C}$ ) for IVCON(i) = 1. Conductivity $\left( \frac{\text{W}}{\text{m} \cdot \text{K}} \right)$ at liquid saturation = 1 for IVCON(i) = 2.
VC2F(i)	real	Reference conductivity $\left( \frac{\text{W}}{\text{m} \cdot \text{K}} \right)$ for IVCON(i) = 1. Conductivity $\left( \frac{\text{W}}{\text{m} \cdot \text{K}} \right)$ at liquid saturation = 0 for IVCON(i) = 2.
VC3F(i)	real	Change in conductivity with respect to temperature for IVCON(i) = 1. Not used for IVCON(i) = 2.
IVCND	integer	Number referring to the sequence of models read in Group 1. The default is [1].

### 6.2.57 Control statement velo (optional)

The input for this macro is identical to macro **flxo**, except that velocities instead of fluxes are calculated (see page 39).

### 6.2.58 Control statement wlbr (optional)

Not supported in this implementation of FEHM.

### 6.2.59 Control statement zone (optional)

Geometric definition of grid for input parameter assignment. The default is input by nodes.

Group 1 - IZONE

Group 2 - X1, X2, X3, X4 (for 2-D) or X1, X2, X3, X4, X5, X6, X7, X8 (for 3-D)

Group 3 - Y1, Y2, Y3, Y4 (for 2-D) or Y1, Y2, Y3, Y4, Y5, Y6, Y7, Y8 (for 3-D)

Group 4 - Z1, Z2, Z3, Z4, Z5, Z6, Z7, Z8 (for 3-D problems only)

The following alternate form of input may be used (starting with Group 2):

Group 2 - MACRO

Group 3 - XG, YG (for 2-D) or XG, YG, ZG (for 3-D) (used with 'list' option)

or

Group 3 - NIN, NODE(1), . . . , NODE(NIN) (used with 'nnum' option)

The geometric-zone description is implemented by defining geometric regions. The coordinates given in Groups 2, 3, and 4 refer to the node positions shown in Fig. 2. All properties defined by node (JA, JB, JC) in any control statements may be defined by **zone**. In the previous macro descriptions, if  $JA < 0$ , then the zone  $IZONE = \text{ABS}(JA)$  is referenced.

It is a good policy to refer to the input check file to insure that node assignments have been made as expected. When x-, y-, z-coordinates are used to define zones, boundaries of those zones may be slightly different than specified. This effect is due to the inclusion of volume from elements adjoining included nodes.

When macro statements **dmdp** and **dual** are used, additional zone definitions are automatically generated. These are identified by zones 101-200 for the first set of matrix nodes and 201-300 for the second set of matrix nodes. For example, Zone 101 corresponds to the matrix material occupying the same space occupied by the fracture identified by Zone 1. Furthermore, Zone 201 refers to the second matrix layer in the **dual** control statement.

The macro **zone** must precede the usage of a ZONE reference. **zone** is ended with a blank line. **zone** can be called more than once and regions redefined. When this is done, all previous zone definitions are eliminated. A node may be included in only a single zone at a time.

Input variable	Format	Description
IZONE	integer	Zone identification number for geometric input.
X1-X8	real	X-coordinates defining zone IZONE.
Y1-Y8	real	Y-coordinates defining zone IZONE.
Z1-Z8	real	Z-coordinates defining zone IZONE.

Input variable	Format	Description
MACRO	character*4	String denoting alternate input format. MACRO = "list" - read a list of x-, y-, z-coordinates, one set per line until a blank line is encountered. The nodes corresponding to these coordinates make up the zone. MACRO = "nnum" - read the number of specified nodes, followed by the node numbers. These comprise the zone.
XG	real	X-coordinate of node to be included in IZONE.
YG	real	Y-coordinate of node to be included in IZONE.
ZG	real	Z-coordinate of node to be included in IZONE.
NIN	integer	Number of nodes in IZONE.
NODE(i)	integer	NIN node numbers of the nodes to be included in IZONE.

The following is an example of **zone**:

zone			
1			
0.00	1000.00	1000.00	0.00
1075.00	1074.00	1079.00	1080.00
2			
0.000	1000.000	1000.00	0.00
870.000	869.000	1074.00	1075.00
3			
0.000	1000.000	1000.00	0.000
860.000	859.000	869.000	870.000
4			
0.000	1000.000	1000.00	0.000
780.000	779.000	859.000	860.000
5			
0.000	10000.00	1000.00	0.000
730.000	730.000	779.000	780.000
6			
0.000	1000.000	1000.00	0.000
700.000	700.000	730.000	730.000

## 7.0 OUTPUT

Output is found in the code-generated files (output file, write file, history plot file, solute plot file, contour plot file, contour plot file for dual or dpdp, stiffness matrix data, input check file, and AVS output files) described in Section 5.0.

Macro commands (input options) dealing with output control are **cont** (page 27), **ctrl** (page 30), **nod2** (page 46), **node** (page 46), and **time** (page 66). The command **cont** is used to specify output format and time intervals for contour data output (*fehm.con*, *fehm.dp*); **ctrl** is used to specify if element coefficients calculated in the code should be saved (*fehm.stor*); **node** and **nod2** are used to provide nodal or coordinate positions for which general information and time history data will be output (*fehm.out*, *fehm.his*, *fehm.trc*, and terminal output); and **time** provides input on the time printout interval for nodal information (*fehm.out* and terminal output).

The code itself provides no graphical capabilities. A software environment, called the Browser, provides preprocessing, processing, and postprocessing capabilities and a help-index for FEHM. A complete description of how to use the Browser is given in Section 8.5.1. The Browser Pre-Processor assists the user in setting up an input file by displaying the appropriate pages from this User's Manual, doing syntax checks, analyzing the input file, and providing direct access to auxiliary programs for structured and unstructured grid generation.

The Browser Processor allows the user to run FEHM, restart it, and provides assistance in setting up a control file.

The Browser Post-Processor provides FEHM time history plots, contour plots, and access to AVS (an AVS license is required for its use).

Time history plots of the energy source, source strength, temperature, pressure, capillary pressure, and saturation are made from the *flen.his* FEHM output files. Data from the *flen.trc* files are used to make time history tracer plots of the 10 species concentrations. Capabilities for zooming, scaling, using a log scale, and printing the data are provided.

Contour plots using 2-D quad grids and 3-D hex grids for material properties, temperature, saturation, pressure, velocities, and solute concentrations can be made from the AVS FEHM output files. The plots can be rotated, zoomed, scaled, translated, and printed. Axis values and the color bar can be customized.

AVS provides tools for visualization and analysis of volumetric scalar and vector data. AVS FEHM output files are available for the following node data: material properties, liquid- and vapor-phase values, velocity and pressure values, temperature, saturation, concentration dual, and dpdp. The AVS output files from FEHM are written in either an ASCII or binary format that can be imported into AVS UCD graphics routines for viewing.

Two additional graphical postprocessing routines are available for use with output files *flen.his*, *flen.trc*, and *flen.con*, as discussed in Section IIIC of Zyvoloski et al. (1991).

Additional information on the data found in the output files is given below.

## 7.1 Output File (*filen.out*)

Information contained in the general output file is mostly self explanatory. The file starts with the code version, date, and time followed by the user input problem title. A summary of the I/O files used, macro control statements read, and array storage follow. Variable information for user-specified nodes at user-selected time intervals is written. The file ends with a summary of simulation time, number of time steps in the problem, the number of iterations taken, and total cpu time.

## 7.2 Write File (*filen.fin*)

The write file contains the final values of pressure, temperature, saturation, and simulation time for the run. The final version of the file is generally written at the end of the simulation. This information is also written if the minimum user-supplied time step has been realized and execution is terminated. If the write file has not been specified at startup, the code will use *fehmn.fin*. The primary use of the write file is as a restart file. The write file contains the following:

- Code version number, date, time

- Problem title

- Simulation time (days)

- Gas flag (ngas, h20, air)

- Tracer flag (trac, ptrk, ntra)

- Stress flag (strs, nstr)

- Dpdp flag (dpdp, ndpd)

- Dual flag (dual, ndua)

If ngas flag is set, followed by:

- Final temperature (°C) at each node

- Final saturation (dimensionless) at each node

- Final pressure (MPa) at each node

- Final capillary pressure (MPa) at each node

Or if neither the air or ngas (h20) flag are set, followed by:

- Final temperature (°C) at each node

- Final saturation (dimensionless) at each node

- Final pressure (MPa) at each node

Or if air flag is set, followed by:

- Final saturation (dimensionless) at each node

- Final pressure (MPa) at each node

If trac flag is set, followed by:

Number of species

Species concentration (dimensionless) for each node for each species

Or if ptrk flag is set, followed by:

Number of particles, final random number seed

Final node position for each particle (if the value is negative, the particle left the model domain at a fluid sink at that node)

Fractional time remaining at current node for each particle

Multiplier to the plug flow residence time for each particle at the current node position, accounting for dispersion, sorption, and matrix diffusion effects

Age for each particle, i.e., the time since the particle entered the system. However, if the particle has left the system, this value is the time the particle left.

If the random number seed in the file is negative, the arrays for the fractional time remaining and the multiplier to the plug flow time have been omitted using the PRNT\_RST = -1 option (see PRNT\_RST description in the PTRK macro). A restart simulation using this input file will only approximate the behavior of particles because each particle will be assumed to have just entered the node. It is preferable to restart a particle-tracking simulation using a file that contains the full restart information.

If strs (not implemented in this version)

If dpdp or dual flag was set:

The above information including dual-porosity/dual-permeability nodes

### 7.3 History Plot File (*flen.his*)

The history plot file contains the following:

Code version number, date, time

Problem title

Tracer flag ('trac' or blank)

Stress flag ('strs' or blank)

Number of nodes for which data are output

Node number and x-, y-, and z-coordinate (m) of each node for which data are output

'headings'

'node flow enthalpy (Mj/kg) flow (kg/s) temperature (deg C) total pressure (Mpa)'



For ngas:

'air pressure (Mpa) capillary pressure (Mpa) saturation (kg/kg) relative humidity'

For airwater:

'capillary pressure (MPa) saturation (kg/kg)'

And for each time step:

Time (days) followed by:

For ngas:

Node number, energy source (MJ/s), source strength (kg/s), temperature (°C), total pressure (MPa), air pressure (MPa), capillary pressure (MPa), saturation (dimensionless), and relative humidity for each specified output node

For airwater:

Node number, energy source (MJ/s), source strength (kg/s), temperature (°C), pressure (MPa), capillary pressure (MPa), saturation (dimensionless) for each specified output node

## 7.4 Solute Plot File (*filen.trc*)

Solute data are output for the same nodes used for the history plot file. The solute plot file contains:

Code version number, date, time

Problem title

Number of nodes for which data are output.

Node number and x-, y-, and z-coordinate (m) of each node for which data are output

Number of different species for tracer solution

And for each time step:

Time (days), species number followed by

Species concentration (dimensionless) for each specified output node.

When particle tracking is used, the concentration can be output in several different forms (number of particles, number per fluid mass, or number per total volume). The choice of which form to use is input in the **ptrk** macro.

## 7.5 Contour Plot File (*filen.con*)

The contour plot file contains:

Code version number, date, time

Problem title

Tracer ('trac') solution or blank

Stress ('strs') solution or blank

Number of nodes for which data are output

X-, y-, and z-coordinate (m) of each node for which data are output

Number of nodes per element, total number of elements

Nodal connectivity information for each node of each element

X-, y-, and z-permeability ( $\text{m}^2$ ) for each node

X-, y-, and z-thermal conductivity ( $\frac{\text{W}}{\text{m} \cdot \text{K}}$ ) for each node

Porosity, rock specific heat ( $\frac{\text{MJ}}{\text{kg} \cdot \text{K}}$ ), capillary pressure (MPa) for each node

Number of degrees of freedom per node for the current problem, direction of gravity in problem, value of gravity

If tracer solution is present:

Number of species

And for each specified time:

Time (days), injection phase ( $\geq 0$  liquid,  $< 0$  vapor) followed by

If injection phase is liquid:

Liquid transmissibility/density, liquid density ( $\text{kg}/\text{m}^3$ ), pressure - capillary pressure (MPa), temperature ( $^{\circ}\text{C}$ )

And if tracer solution is present:

Species concentration of liquid phase

Or if injection phase is vapor:

Vapor transmissibility/density, vapor density ( $\text{kg}/\text{m}^3$ ), pressure (MPa), temperature ( $^{\circ}\text{C}$ )

And if tracer solution is present:

Species concentration of vapor phase.

## 7.6 Contour Plot File for dual or dpdp (*filen.dp*)

The contour plot file for dual or dpdp contains the same information as the regular contour plot file only the parameter values are for the dual-porosity/dual-permeability nodes.

## 7.7 Stiffness Matrix Data (*filen.stor*)

The stiffness matrix data file is used to store the finite-element coefficients for each node. The file eliminates the need for the code to recompute the coefficients for subsequent runs. It contains the following:

- Code version number, date, time

- Problem title

- Number of storage locations needed to store geometric input types, number of nodes, size of connectivity array

- Volume associated with each node

- Nodal connectivity information for each connection

- Position of geometric coefficient for each connection

- Diagonal position in connectivity array for each node

- Finite-element geometric coefficient for each for each storage location

If stress solution is enabled:

- Finite-element geometric coefficient for each for each storage location for the stress module

## 7.8 Input Check File (*filen.chk*)

This file contains a summary of input information that may be of use in debugging or memory management of the code. The positions of maximum and minimum values of input parameters and derived quantities are given. Also provided is an analysis of array storage requirements.

## 7.9 Error Output File (*fehmn.err*)

This file contains the code version number, date, and time followed by any error or warning messages issued by the code during a run.

## 7.10 AVS Log Output File (*filen.10001\_avs\_log*)

The AVS log output file contains:

- Code version number, date

- AVS log identifier

- Problem title

And for each specified time:

- AVS output file prefix, call number, and time (days)

## 7.11 AVS Header Output Files (*filen.number\_type\_head*)

The data types are given in Section 7.13.1.

### 7.11.1 ASCII header

The AVS ASCII (formatted) header files contain:

20 lines of text with information about the FEHM AVS output files. The text is followed by a one-line AVS UCD file header containing:

- number of nodes
- number of cells
- number of data components for the nodes
- number of data components for the cells (currently 0)
- number of data components for the model (currently 0)

### 7.11.2 Binary header

The AVS binary (unformatted) header files consist of 21 bytes with the following values:

- number '7' indicating binary file (1 byte unsigned char)
- number of nodes (4 byte int)
- number of cells (4 byte int)
- number of node data (4 byte int)
- number of cell data (4 byte int) (currently 0)
- number of model data (4 byte int) (currently 0)

## 7.12 AVS Geometry Output File (*filen.10001\_geo*)

### 7.12.1 ASCII geometry output file

The ASCII (formatted) geometry file contains the following:

- Node id and x-, y-, z-coordinate for each node
- Cell id, material id, cell type, and the list of cell vertices

### 7.12.2 Binary geometry output file

The binary (unformatted) geometry file contains the following:

- Number of nlist nodes (4 byte int)
- Cell id, material id, number of nodes, cell type (16 \* num\_cells) (ints)
- Cell vertice list (4 \* num\_nlist\_nodes) (ints)
- X-coordinates for nodes (num\_nodes \* 4) (floats)
- Y-coordinates for nodes (num\_nodes \* 4) (floats)
- Z-coordinates for nodes (num\_nodes \* 4) (floats)

## 7.13 AVS Data Output Files (*filen.number\_type\_node*)

### 7.13.1 ASCII node data output files

All the ASCII (formatted) node data files contain the following:

Number of data components and size of each component

A label/unit string for each data component

And for each node:

The associated node data (described by data type below):

#### 7.13.1.1 Material properties (*\_mat* and *\_mat\_dual*)

These data will consist of 11 fields. The order of the fields are:

Permeability in x-, y-, and z-direction ( $\text{m}^2$ )

Thermal conductivity in x-, y-, and z-direction ( $\frac{\text{W}}{\text{m} \cdot \text{K}}$ )

Porosity

Rock specific heat ( $\frac{\text{MJ}}{\text{kg} \cdot \text{K}}$ )

Capillary pressure (MPa)

Relative permeability model

Capillary pressure model

The dual or dpdp values for each of these fields will be written to a file with "mat\_dual\_node" appended to the file name.

#### 7.13.1.2 Scalar parameters (*\_sca* and *\_sca\_dual*)

These data files will contain scalar data including:

Saturation

Temperature ( $^{\circ}\text{C}$ )

Liquid pressure(MPa)

Vapor pressure(MPa)

If dual values are calculated, they can be written to the sca\_dual output file.

#### 7.13.1.3 Vector parameters (*\_vec* and *\_vec\_dual*)

These data files contain the vector values for:

Liquid and vapor velocities (m/s)

If `idualp` is defined, dual-porosity values for the vapor phase will be written to the `_vec_dual` file. If `idpdp` is defined, double-porosity/double-permeability values for the liquid phase will be written.

#### 7.13.1.4 Solute concentrations (`_con` and `_con_dual`)

Up to 20 fields per node can be written for solute concentrations. The number written is determined by the number of species. The dual counterparts to each will be written to the `_con_dual` file.

### 7.13.2 Binary node data output file

All the binary (unformatted) AVS data files contain the following:

Node data labels (1024 byte string)

Node data units (1024 byte string)

Number of node components (4 byte int)

Node component list (`num_node_data * 4`) (ints)

Minimums for node data (`num_node_data * 4`) (floats)

Maximums for node data (`num_node_data * 4`) (floats)

Data blocks with values for each node ( `num_nodes * num_node_data * 4`) (floats)

The data types are described above in Section 7.13.1.

## **8.0 SYSTEM INTERFACE**

### **8.1 System-dependent Features**

In addition to standard intrinsic math routines, only two system routines are required by the FEHM code. The code uses a system call to get the date (subroutine dated) and a system routine to get the CPU clock time (subroutine tyming).

### **8.2 Compiler Requirements**

FEHM is written in Fortran 77 and C. FEHM has been successfully compiled and run on SUN, HP, IBM RISC, SGI, and Cray computers.

### **8.3 Hardware Requirements**

No special hardware features or environments are required by the software.

### **8.4 Control Sequences or Command Files**

None.

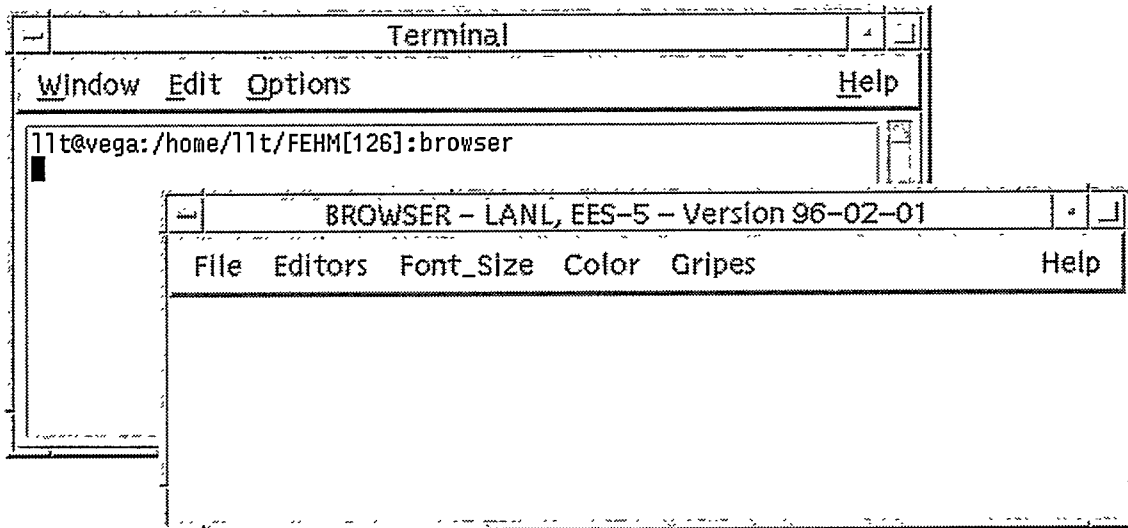
### **8.5 Software Environment**

FEHM can be run under a software environment called the Browser. The Browser is a column-oriented graphical user interface that provides preprocessing, processing, and postprocessing capabilities and a help index for FEHM.

#### **8.5.1 Browser**

The Browser Pre-Processor assists the user in setting up an input file to be run with the Processor and provides direct access to auxiliary programs for grid generation. The Processor allows the user to run FEHM and restart it. The Post-Processor allows the user to view ASCII output files, make time history plots, make contour plots, and display AVS output files generated by FEHM (user must have an AVS license). There is also a help index that includes documentation for FEHM and auxiliary programs. To bring up the main Control Window for the Browser under X-windows, type:

**browser**



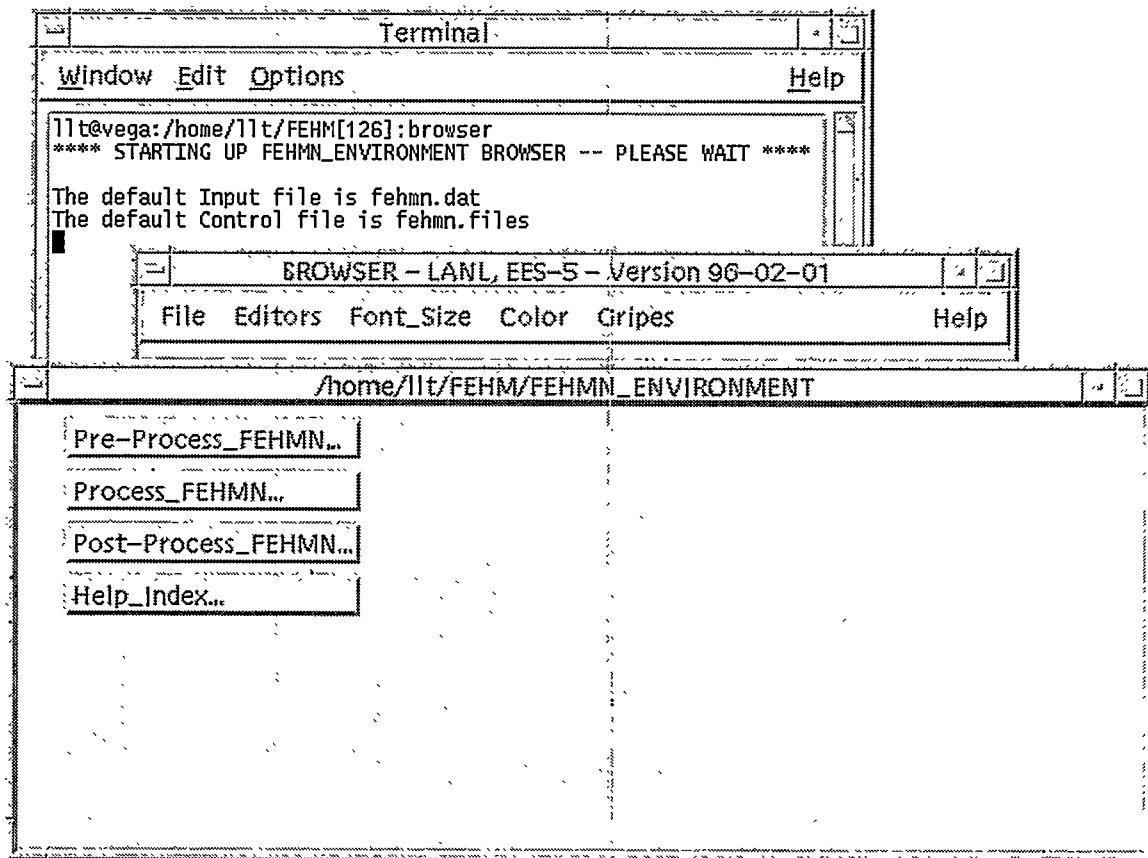
- Control Window

The *Editors* button along the top allows the user to switch editors. The default editor is *vi*. The editors *emacs* and *fred* are menu options for the Suns. *Jot* is a menu option for the SGIs. The *Font\_Size* button allows the user to change the font size. A medium-size font is the default. The *Color* button allows the user to use the Browser on a black-and-white terminal. The *Gripes* button allows the user to report bugs and view a list of current bugs.

The *Help* button has a *Getting\_Started* menu option that gives more details on using the above buttons. The *Help* button also allows the user to turn the FEHM User's Manual pages on or off. The default is to have the manual pages on.

To bring up the Browser window, select *Open* in the Control Window under *File*, highlight the file *FEHMN\_ENVIRONMENT*, and click on the *OK* button. The following Browser Window will appear.





- Browser Window

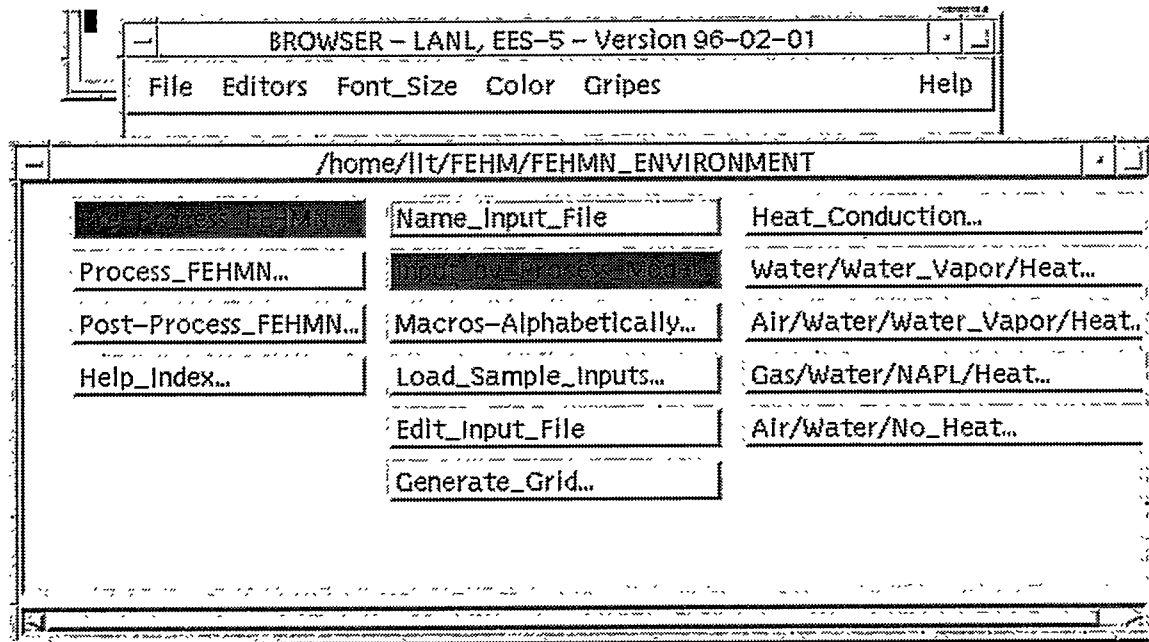
Use the Browser by clicking ONCE with the LEFT mouse button on the desired button. If the button name is suffixed with "...", another column of buttons will appear when selected. Otherwise, a function will be executed. Red buttons are the last buttons selected. Green buttons were pushed previously. Grey buttons have not been accessed.

The scroll bar can be used to reach buttons outside the displayed window, or the window can be resized by dragging the corner of the window.

When finished, quit the Browser from the Control Window, under *File*, by selecting *Exit*.

#### 8.5.1.1 Pre-Process FEHM

The *Pre-Process\_FEHMN* button provides guidance, syntax checking, analysis, and help in creating an input file. Several auxiliary grid-generating programs are provided. Click on *Pre-Process\_FEHMN* to get the column with these options.



- Create Input File

The default input file name is *fehmn.dat*. To change this, the user should click on the button *Name\_Input\_File*. This file name must be assigned before working on the file. There are four ways the user can work on this file. A user can input by process model type, from a list of macros, by loading a sample input file, or by editing the input file directly. A combination of these ways can be used.

The *Input\_by\_Process\_Model* button allows the user to select the type of problem from five process models and guides the user through the required and optional macros for that process type. In the example above, the *Heat\_Conduction* process model was selected. The user clicks on the *Required\_\*\_Inputs* button to get a list of the macros required for a specific process type. When a button is clicked for a specific macro, the appropriate pages from the FEHM User's Manual are displayed and an editing window is opened for the user to input the macro. The input of macros are appended to the input file. A macro may be written to this file more than once. Displaying the FEHM User's Manual pages can be turned off under *Help* on the main Control Window. The editor can be switched by clicking on the *Editors* button in the main Control Window. As input is entered, a syntax check is made for the correct number and type of arguments. Corrections are made, if possible, and displayed. The *Analyze\_Input* button can be used to verify that all the required macros are present and to list the appropriate optional macros that have not been included. There is an option, the *Use\_Sample\_File\_\** button, for using an example

input file for each process type. The example input file is loaded into this file. The *Edit\_Input\_File* button gives the user direct access to this input file at any time.

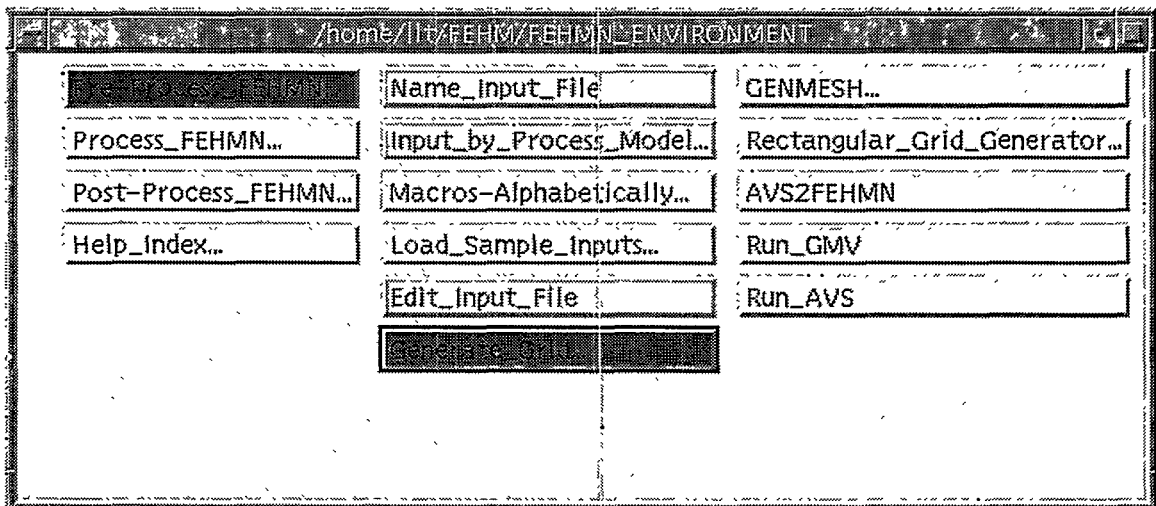
The *Macros-Alphabetically* button provides an alphabetical listing of all the macros. This mode will automatically bring up the FEHM User's Manual pages and an editing window. Macros entered are appended to the input file, and a syntax check is made on the input. Direct access to the input file is also available through the *Edit\_Input\_File* button. There is no input guidance or analysis of required and optional macros.

The *Load\_Sample\_Inputs* button replaces or creates a new input file. Direct access to the input file is available by going through the *Edit\_Input\_File* button. Access to the User's Manual is available through the *Help\_Index* in the first column.

The *Edit\_Input\_File* button allows the user to directly edit the input file. The editor can be changed with the *Editors* button in the Control Window. The User's Manual is available by going through the *Help\_Index* in the first column. There is no syntax checking, input guidance, or analysis on the input.

- Generate Grid

Direct access is given to several auxiliary grid-generating programs. A brief description is given below. Additional help is available under the *Help\_Index* button.



*GENMESH* is an automatic mesh-generation code that directly generates the finite-element mesh input required by FEHM. Help pages and an editing window are used to create the input file. Examples of input files for 2-D and 3-D grids

are provided. As mentioned earlier, the User's Manual is available in the Browser.

*Rectangular\_Grid\_Generator* program creates a structured grid in the format for an FEHM input file. If the file *input.grid* is constructed, it can be run with an input file. Otherwise, an interactive mode is available that prompts for the necessary inputs.

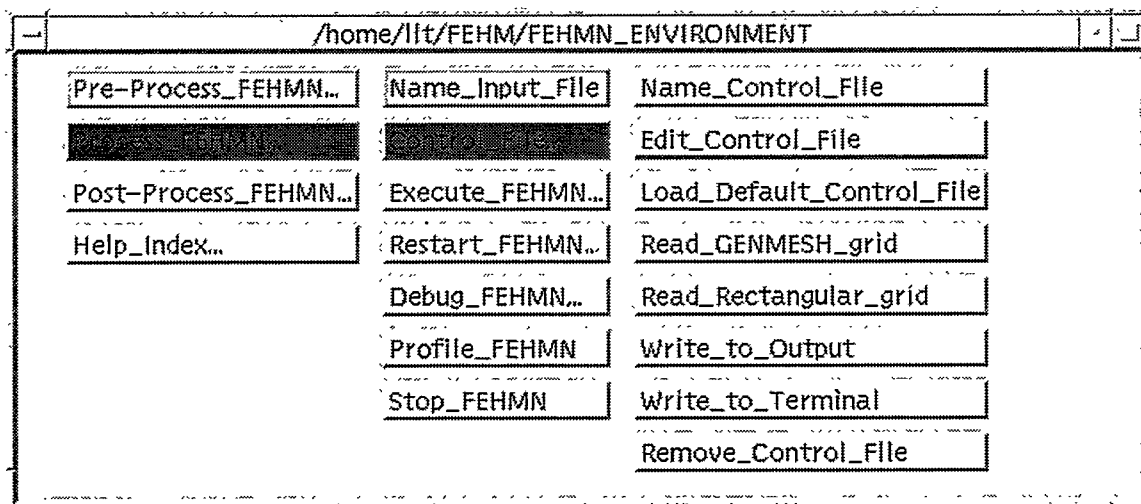
*AVS2FEHMN* takes an AVS file and converts it to the format for an FEHM input file.

*Run\_GMV* is a General Mesh Viewer, a visualization tool that provides 3-D interactive graphics for data from any 3-D mesh.

*Run\_AVS* executes the commercial program AVS. (An AVS license is required.)

### 8.5.1.2 Process FEHM

The *Process\_FEHMN* button provides assistance in creating a control file and in executing, restarting, and stopping FEHM. The input file name will be the name used under *Pre-Process\_FEHMN*, or it can be changed with the *Name\_Input\_File* button under *Process\_FEHMN*.



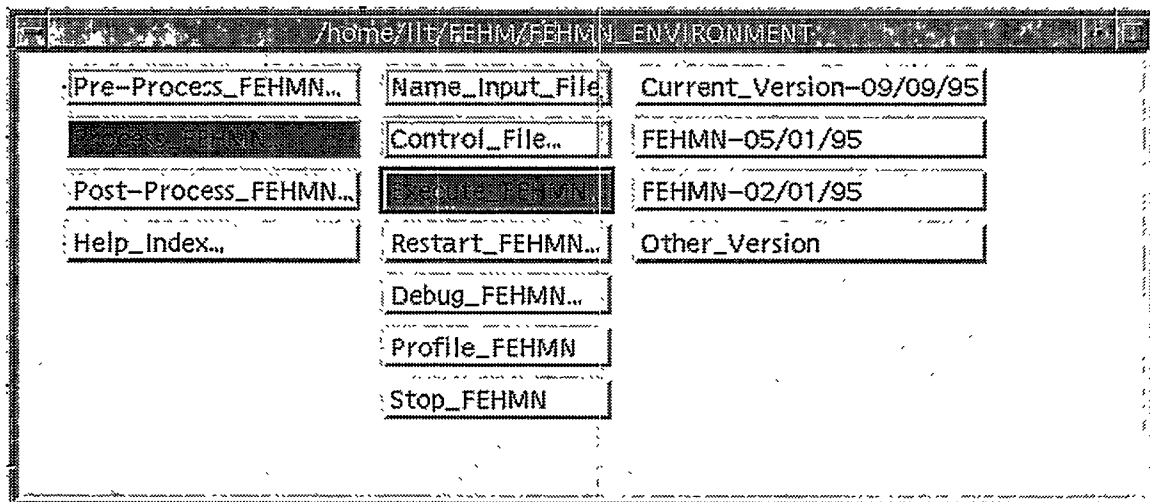
- Control Files

The *Control\_File* button allows users to create their own control file or have the Browser create the appropriate control file. The default control file name is *fehmn.files*. Another name can be entered with the *Name\_Control\_File* button. The *Edit\_Control\_File* button allows the user to directly edit the control file. The *Load\_Default\_Control\_File* button will create a control file based on the name of the current input file. The *Read\_GENMESH\_grid* and *Read\_Rectangular\_grid* buttons modify (or create, if none exists) the control file to read the appropriate grid input file. The

*Write\_to\_Output* button modifies (or creates, if none exists) the control file to write the output to a file. The *Write\_to\_Terminal* button modifies (or creates if none exists) the control file to write the output to the terminal. If you do not want FEHM to automatically run with a control file, *fehmn.files* can be deleted with the *Remove\_Control\_File* button.

- Executing, Restarting, Stopping FEHM

Options for the most current version of FEHM and older, frozen versions are available under the *Execute\_FEHMN* button. Simply click on the version needed. If the control file *fehmn.files* is in your local space, the program will automatically begin. If another control file name is used, an interactive mode will start in an xterm window and the control file name can be entered at the first prompt. If no control file is used, the user will be prompted to input file names.



Options for the most current version of FEHM and older, frozen versions are available to continue a run under the *Restart\_FEHMN* button. When these are clicked on, a help page will pop up that describes how the macro *time* needs to be adjusted for a restart run. The control file will be changed for a restart run, or if there is no control file, one will be made. The *\*.fin* file will be copied to *\*.ini*.

Options for the most current version of FEHM and older, frozen versions are also available for a debug mode and a profile mode under the *Debug\_FEHMN* and *Profile\_FEHMN* buttons, respectfully.

FEHM can be stopped at any time by using the *Stop\_FEHMN* button. This button will give you the process id and directions on how to kill the process in the xterm window provided.

### 8.5.1.3 Browser FEHM Post-Processor

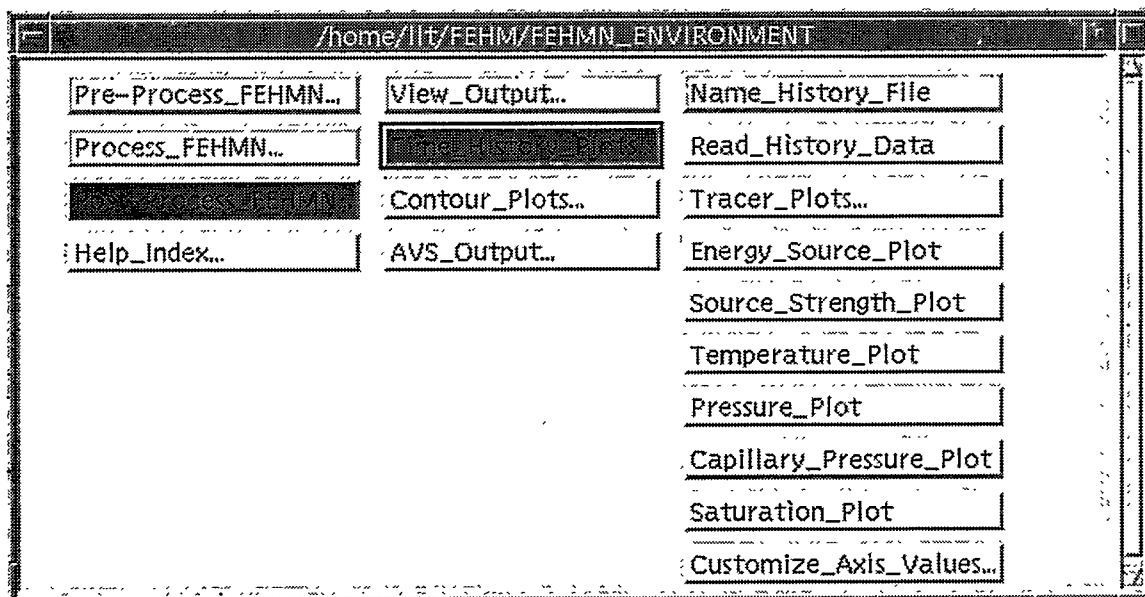
The Browser Post-Processor allows the user to view FEHM output files, make time history plots, make contour plots, and display the output with AVS (an AVS license is required).

- Viewing FEHM Output Files

The *View\_Output* button will allow the user to view the FEHM output files. The name used for the \* (wildcard character) will be the prefix of the last input file name used or the name entered in the *Name\_Output\_File* button.

- Time History Plots

The *Time\_History\_Plots* button allows the user to make time



history plots of the energy source, source strength, temperature, pressure, capillary pressure, and saturation from the FEHM \*.his and time history Tracer plots of the 10 species concentrations from FEHM \*.trc output files. \*.his is specified on the seventh line of the control file, and \*.trc on the eighth line (see Section 6.2.1 for more information on how to set up your control file to create these output files). A line for each node specified in the input is plotted as time versus the value. The user can zoom in on the plots, resize the window, and use a log scale for the x- or y-axis.

The title on the first line of the input deck is used for the title of the plot. The second title line displays the version of FEHM and the date run. The x and y axes are automatically scaled. A legend is automatically displayed showing which line corresponds to which node.

The default time history file is *fehm.his*. To change this, the user should click on the button *Name\_History\_File*. The data will automatically be converted to xrt plotting format when a new history file is named. To read in new data with the same file name (or if using the default), select the *Read\_History\_Data* button. This action will convert the new data to the xrt plotting format.

Once the data are converted, plots can automatically be brought up by selecting the button for the plot desired. The *Tracer\_Plots* button will bring up another column listing the 10 species. Only the species in the \*.trc file will be available for plotting. A message will be given if a species is selected that was not in the \*.trc file.

The values for the axis can be specified. Select the *Customize\_Axis\_Values* button, the *New\_XY\_Values* button, then the axis you want to specify. You can get the minimum and maximum values or return to having it automatically specified with the *Default\_XY\_Values* button. After setting the axis values, select the *Save\_Axis\_Values* button to save them.

Following are the controls available to manipulate the time history plots:

**Zooming:** Press control and hold down the left mouse button.  
Move the mouse to draw a box around the area to zoom into.

**Scaling:** Press control and hold down the middle mouse button.  
Move the mouse down to increase the graph's size.  
Move the mouse up to decrease the graph's size.

**Reset:** To reset the window, type r.

**Resizing Window:** The window size can be changed by clicking on a corner of the window and dragging the mouse to the desired size.

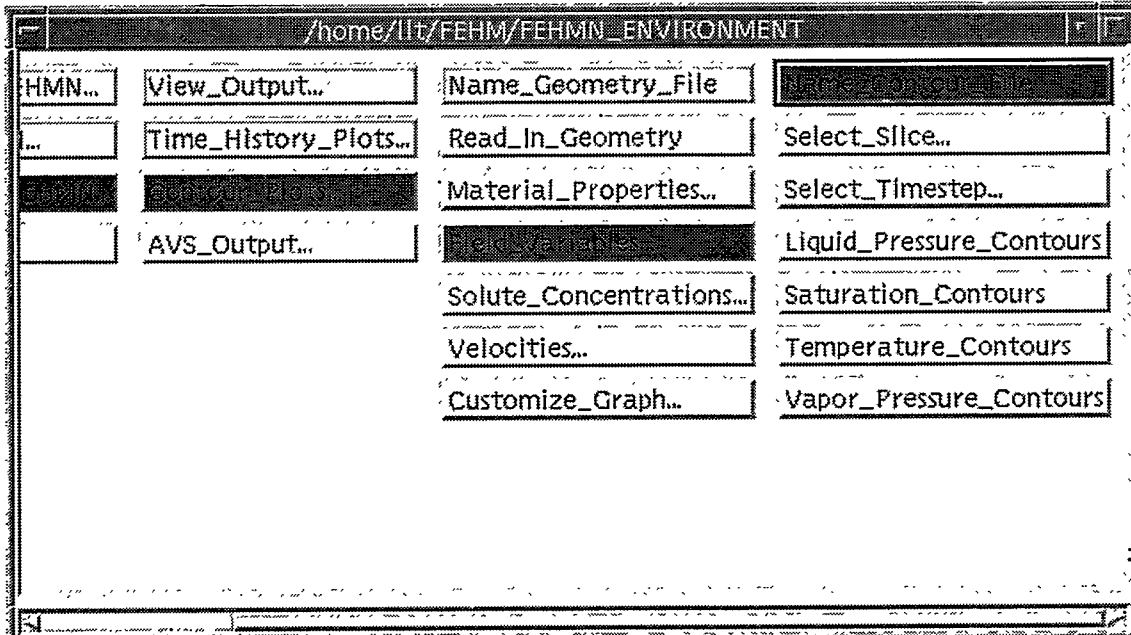
**Log Scale:** The x- and y-axes can each be changed to a log scale by clicking on the toggle button on the top row.

**Printing:** Click on the *Print* button on the top row. The default is set for the printer named *graphics*. This default can be changed by editing the name. A postscript file or a XWD file can also be printed. Other options can be changed under the *Properties* button along the bottom.

Exiting: Click on the *Exit* button on the top row.

- Contour Plots

The *Contour\_Plots* button allows the user to make plots of x, y, or z slices of material properties, temperature, saturation, pressure, velocities, and solute concentrations for 2-D quad and 3-D hex grids. The plots can be rotated freely in the x-, y-, or z-direction or from any end point. The plots can be zoomed, scaled, translated, and the window can be resized.



There are options for drawing the mesh, drawing shaded surfaces, drawing contour lines, and drawing zones.

The title on the first line of the input deck is used for the title of the plot. The version of the FEHM executable and date run is the second line of the title. A title at the bottom of the plot includes the type of plot, the slice taken, and which time step, if applicable. The x-, y-, and z-axis and the color bar are automatically scaled, unless specific values are given. A legend is automatically displayed showing the numeric value of the colors.

The contour plots use data from FEHM AVS output files created by specifying the **cont** macro with **avs** option in the FEHM input file (see Section 6.2.8). Slices of material-property contour plots can be made by specifying **material** after **avs**. The contour plots include permeability in the x-, y-, or z-direction; thermal conductivity in the x-, y-, or z-direction;



porosity; rock specific heat; capillary pressure; and models for relative permeability and capillary pressure.

Time-dependent slices of field-variables plots can be made for liquid pressure, vapor pressure, saturation, and temperature. A combination of **pressure** and **liquid or vapor, saturation, temperature** after **avs** in the **cont** macro is needed.

Time-dependent slices of solute concentrations can be made by specifying **concentration** after **avs** in the **cont** macro. Buttons for the 10 species are provided.

Time-dependent slice of velocities can be made by specifying **velocity** and **liquid or vapor** after **avs** in the **cont** macro. The speed (the square root of the sum of the squares of the x, y, and z velocities) for liquid and vapor can be displayed.

The default AVS geometry file is *fehmn.10001\_geo*. To change this, the user should click on the button *Name\_Geometry\_File*. The geometry will automatically be converted to xrt plotting format when a new geometry file is named. To read in new data with the same geometry file name (or if using the default), select *Read\_In\_Geometry*. This action will convert the new data to the xrt plotting format. The same geometry can be used with multiple contour files.

The default AVS contour files are *fehmn.1000\**. To change this, the user should click on the button *Name\_Contour\_File*. The contour data will be read from the file selected.

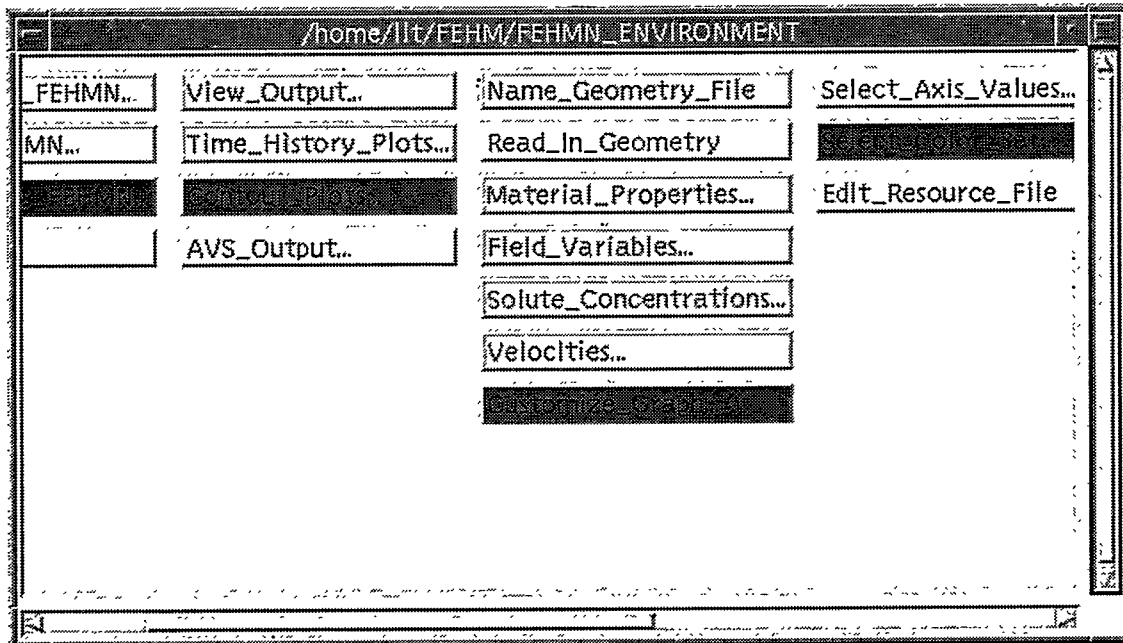
Once the geometry is converted and a contour file named, plots can automatically be brought up. For material-property plots with 3-D grids, the user must select the type of slice and the value for the slice. If not selected, the default will be an xy slice with a z-value of 1.0. The geometry values (from the file *\*.10001\_geo*) can be displayed by clicking on the button *Display\_Geometry\_File*. To display the plot, the user should click on the plot type. For 3-D grids, the slice type and value will remain the same for additional plot types until changed.

For field-variable, solute-concentration, and velocity plots with 3-D grids, the user must select the type of slice and the value for the slice. If not selected, the default will be an xy slice with a z-value of 1.0. For 2-D and 3-D grids, the user must also select the time step (the default is the second time step). The applicable time steps (from the file *\*.10001\_avs\_log*) can be viewed by clicking on the button *Display\_Time\_Steps*. The

user can enter the time step desired with the *Input\_Time\_Step* button or the value of the day desired with the *Input\_Number\_Days* button. To display the plot, the user should click on the plot type. The slice type, value, and time step will remain the same for additional plot types, until changed.

The values for the axis can be specified. Select the *Customize\_Graph* button, then the *Select\_Axis\_Values* button and the axis you want to specify. You can get the minimum and maximum values or return to having it automatically specified.

The number of colors and the colors can be specified. Select the *Select\_Color\_Bar* button. The colors for the color bar can be selected from a range of colors with the *Select\_Colors* buttons, and the number of color divisions can be selected with the *Change\_Number\_Color\_Divisions* button.



Other available color buttons are *Use\_Standard\_Color\_Bar*, *Use\_Random\_Colors*, *Use\_Gray\_Scale*, *Use\_No\_Color* (black & white). To select any color available on the system, use the *Display\_Available\_Colors* button; to return to having automatically generated color bars, use the *Use\_Auto\_Color\_Bar* button. The files containing the list of colors and the number of color divisions can also be directly edited with the *Edit\_Color\_List* button and the *Edit\_Color\_Divisions*. Colors are appended to the color list when the *Select\_Color* buttons are selected. To pick up new colors, click on the *Edit\_Color\_List* button and delete the list contents of this file first.

With the *Edit\_Resource\_File* button, the graph and axis titles can be modified. The rotation of the graph can also be modified.

Following are the controls available to manipulate the contour plots:

**Zooming:** Press control and hold down the left mouse button. Move the mouse to draw a box around the area to zoom into.

**Scaling:** Press control and hold down the middle mouse button. Move the mouse down to increase the graph's size. Move the mouse up to decrease the graph's size.

**Translation:** Press shift and hold down the middle mouse button. Move mouse to shift the graph.

**Reset:** To reset the window (for zooming, scaling, and translation only), type r.

**Rotating:** Hold down the middle mouse button and either:

- move mouse counter-clockwise to rotate view clockwise or

- press x, y, z, or e to select an axis and then move mouse perpendicular to axis.

**Resizing Window:** The window size can be changed by clicking on a corner of the window and dragging the mouse to the desired size.

**Printing:** Click on the *Print* button, on the top row. The graph can be printed to any printer or a file. It can create postscript, XWD, or CGM files. Other options can be changed under the *Properties* button along the bottom.

**Exiting:** Click on the *Exit* button on the top row.

Various displays can be made using combinations of the four Toggles at the top of the display:

**DrawMesh:** When highlighted, displays the x-y grid projected onto the 3-D surface in a 3-D view with a z-axis. The graph honors rotation and perspective control.

**DrawShaded:** When highlighted, displays the data as a flat shaded surface in a 3-D view with a z-axis. The graph honors rotation and perspective control.

**DrawContours:** When highlighted, examines the distribution of the data and draws contour lines demarcating each of the distribution levels.

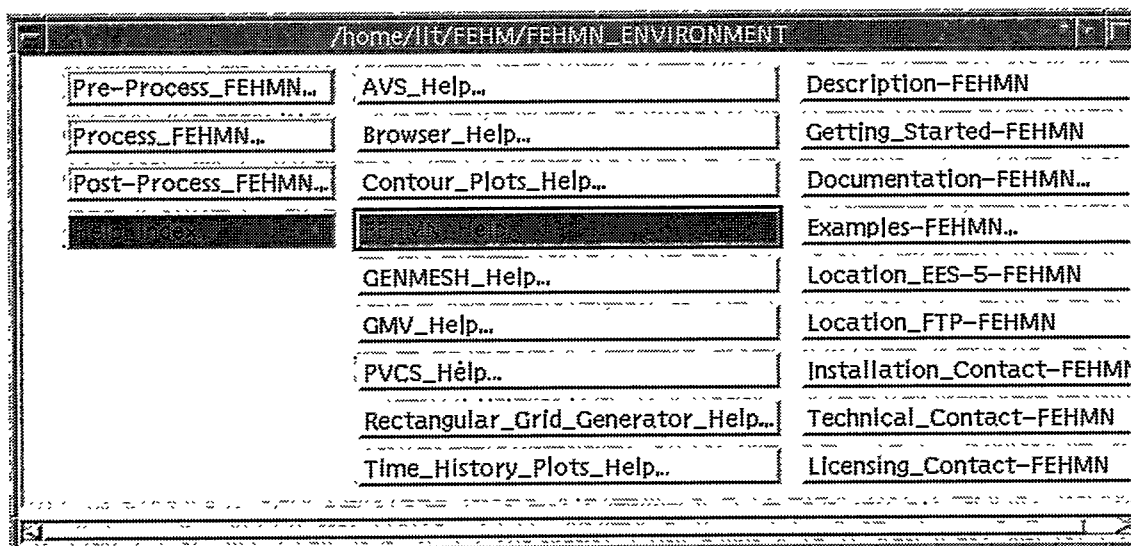
**DrawZones:** When highlighted, examines the distribution of the data and fills each level with a solid color.

- **Display AVS Output**

The *Run\_AVS* button will execute the commercial program AVS (an AVS license is required). The name of the FEHM AVS output file is entered within AVS.

#### 8.5.1.4 Browser help index

The *Help\_Index* button allows the user to view the FEHM documentation and documentation for other codes used within the Browser.



## 8.6 Installation Instructions

On the EES-5 computer network, no installation is required to run FEHM. Table VI lists the location and executable name for each platform.

FEHM object code is provided for users modifying a few routines to link against for installation. FEHM source code is provided for users needing the entire source and for remote users.

Remote users can obtain FEHM executable through a passworded ftp account after a software license agreement form has been completed.

Table VI. FEHM executable locations		
Platform	Path	Executable name
Solaris	/pvcs.config/fehmn/fehmn_96_05_07/objects_sol	xfehmn
IBM	/pvcs.config/fehmn/fehmn_96_05_07/objects_ibm	xfehmn
HP	/pvcs.config/fehmn/fehmn_96_05_07/objects_hp	xfehmn
SGI	/pvcs.config/fehmn/fehmn_96_05_07/objects_sgi	xfehmn

### 8.6.1 Installation of FEHM using objects

Users that need to modify a few routines but do not need the entire source should only have the source they are changing in their local space. On the EES-5 Sun network, the following makefile should be used:

```
/pvcs.config/fehmn/ref/makefile
-OR-
/pvcs.config/fehmn/ref/makefile-g (for debug)
```

and FEHM is installed by typing:

```
make
-OR-
make -f makefile-g (for debug)
```

On the EES-5 IBM (magma), the following makefile should be used:

```
/pvcs.config/fehmn/ref/makefile_ibm
-OR-
/pvcs.config/fehmn/ref/makefile_ibm-g
```

and FEHM is installed by typing:

```
make -f makefile_ibm
-OR-
make -f makefile_ibm-g (for debug)
```

All makefiles create an executable called:

```
xfehmn
```

### 8.6.2 Complete installation of FEHM

Users that need the entire source and remote users can automatically setup the FEHM directory structure. On the EES-5 Sun network, get the following tar files:

```
/pvcs.config/fehmn/fehmn_current/src_files.tar
```

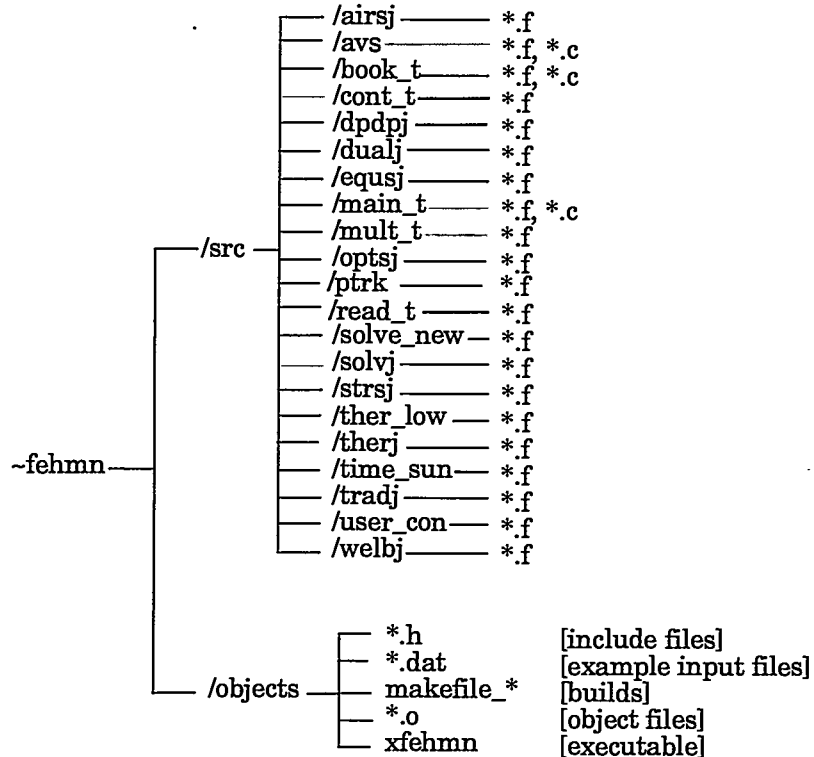
/pvcs.config/fehmn/fehmn\_current/objects\_files.tar

and then type:

```
tar xvf src_files.tar
```

```
tar xvf objects_files.tar
```

The FEHM directory structure should look like the following:



Makefiles are included in these tar files and will be placed in your objects directory. There are makefiles for various platforms with options for debug, pvcs configuration builder commands, and purify. The format is:

```
makefile_machine type[_pvcs or nopvcs][_purify][_g]
```

*Machine\_type* can be *cray*, *hp*, *ibm\_long*, *sgi*, or *sun*. The options inside {} are only available with the *sun* makefiles and are required. The items inside [] are optional. The *purify* option is to link in *purify* and is only available with the *sun* makefiles. The *-g* option is to link in *debug* and is available on all machines. Following are some examples:

makefile\_ibm\_long

makefile\_hp-g

makefile\_sun\_nopvcs

makefile\_sun\_pvcs-g

makefile\_sun\_pvcs\_purify-g

To compile and link FEHM, select the appropriate makefile and type the following:

```
cd ~fehmn/objects  
make -f makefile_sun_nopvcs -OR- make -f makefile_hp
```

All makefiles create an executable called:

```
xfehmn
```

It should be noted that FEHM uses the GZSOLVE application reuse components, solve\_new and slvesu (Zyvoloski and Robinson 1995). Please refer to the SOLVE\_NEW UG component of the document just cited for information on installation of the solver routines.

### **8.6.3 Installation verification and validation**

A series of test scripts have been developed to automate the verification and validation procedure for FEHM. They are described in more detail in the Appendix of "Software Verification and Validation for the FEHM Application" (Dash et al. 1997). Also, see this document for a discussion of the tests performed and their results.

### **8.6.4 FEHM licensing at remote sites**

A software license agreement must be completed before receiving an FEHM executable. To obtain access, get either the Commercial\_agreement.ps or Government\_agreement.ps form from anonymous ftp:

```
ftp ees5-ftp.lanl.gov  
Name (): anonymous  
Password: [enter your email address]  
cd pub/fehm  
ftp> binary  
ftp> get Commerical_agreement.ps
```

or: ftp> get Government\_agreement.ps

The form can be printed with any postscript printer. After reading, please sign the form and return to:

Lynn Trease  
Mail Stop F665  
P.O. Box 1663  
Los Alamos National Laboratory  
Los Alamos, NM 87545  
FAX: 505-665-3687

## 9.0 EXAMPLES AND SAMPLE PROBLEMS

The following describes execution of the FEHM code. Section 9.1 discusses the construction of an input file. Section 9.2 illustrates the entire procedure for executing the FEHM code using terminal input. Example 1 describes the setup and results from a simple 2-D heat-conduction simulation. The remaining sections provide more complex example problems and deal only with problem setup and expected results.

### 9.1 Constructing an Input File

FEHM is a very general simulation code. Thus it is preferable to discuss the construction of an input file from a problem-oriented point of view. In what follows, the needs of the physical problem (initial conditions, boundary conditions, etc.) will be addressed in terms of the macro statements.

**Initial conditions.** These are needed for every problem, even if it is a steady-state simulation. If the simulation is comprised of fully saturated water flow or heat conduction only, then the appropriate control statement would be **init** (page 41). The use of **init** also allows the specification of initial temperature and pressure (gravity) gradients. If two-phase flow is prescribed (thermal or isothermal), then entering the initial conditions through the control statement **pres** (page 49) is more convenient. Initial values for noncondensable gas are handled in the **ngas** (page 44) control statement. It should be remembered that if a restart file is present, those values will have precedence over values input in control statement **init** but not over values input in control statement **pres**. Solute initial conditions are prescribed through the control statement **trac** (page 67).

**Boundary conditions.** Fluid and heat flow boundary conditions can be prescribed through control statements **pres**, **flow** (page 37), **hflx** (page 40), and **rflx** (page 55). Boundary conditions are entered with **pres** by specifying a negative phase-state designation (the code will actually use the absolute value of the phase-state designation). In this case, the code will keep the variable values constant at whatever value was prescribed in **pres**. Flowing pressures are input with the **flow** control statement. Solute boundary conditions are prescribed through the control statement **trac**.

**Material- and energy-balance equations.** The choice of the coupled system equations is made in control statements **sol** (page 64), **ngas**, and **airwater** (page 23).

**Rock or media properties.** These are found in the **rock** (page 58) and **perm** (page 47) control statements.

**Fluid properties.** These are found in control statement **eos** (page 35), which is optional. If **eos** is not invoked, then the properties of water and air included in the code are used. Relative permeabilities, depending on both the fluid and media type, are found in control statement **rlp** (page 55).



**Mesh geometry and nodal coordinates.** This geometry information is found in control statements **coor** (page 29) and **elem** (page 33). This information is usually created with a mesh-generation program.

**Simulation time.** The time-stepping information, including printout intervals and time-step sizing, is found in control statement **time** (page 66).

**Numerics.** Convergence criteria, upwinding parameters, fill-in for the preconditioned conjugate gradient solver and geometry type (2-D, 3-D, radial) are entered with control statement **ctrl** (page 30).

**Advanced iteration control.** Reduced-degree-of-freedom methods are invoked with the **iter** (page 42) control statement. One important quantity entered with this statement is the maximum time for the job to run on the computer.

**Sources and sinks.** These are input with the control statement **flow**. Care must be taken as the parameters have different meanings for different physical models.

Table VII on the next two pages lists the input macros that should be used to formulate various types of problems.

Table VII. Required and optional macros by problem type			
Problem type : Heat conduction		Problem type : Water/water vapor/heat Equivalent continuum, dual porosity*, dual permeability**	
Required macros	Optional macros	Required macros	Optional macros
title	alti	title	alti
cond	cont	cond	cap
coor	finv	coor	cont
ctrl	flo2	ctrl	eos
elem	flxo	elem	finv
flow or hflx	iter	init or pres	flow
init or pres	node or nod2	perm	flo2
rock	renm	rlp	hfxo
sol	rflx	rock	hflx
time	stea	sol	iter
stop	text or comments (#)	time	node or nod2
	user	stop	ppor
	vcon		pres
	zone	dual (* only)	renm
		dpdp (** only)	rflx
			rxn
			stea
			text or comments (#)
			trac
			vcon
			velo
			zone

**Table VII. Required and optional macros by problem type (continued)**

Problem type : Air/water/water vapor/heat or gas/water/NAPL/heat Equivalent continuum, dual porosity* dual permeability**		Problem type : Air/water/no heat Equivalent continuum, dual porosity* dual permeability**	
Required macros	Optional macros	Required macros	Optional macros
title	comments (#)	title	comments (#)
cond	adif	airwater	alti
coor	alti	coor	cap
ctrl	cap	ctrl	cond
elem	cont	elem	cont
flow or hflx	eos	flow or hflx	eos
init or pres	finv	init or pres	finv
ngas	flow	node or nod2	flow
perm	flo2	perm	flo2
rlp	flxo	rock	flxo
rock	hflx	sol	hflx
sol	iter	time	iter
time	node or nod2	stop	ppor
stop	ppor		pres
	renm	dual (*only)	renm
dual (*only)	rflx	dpgp (**only)	rflx
dpgp (**only)	rxn		rxn
	stea		stea
	text		text
	trac		trac
	vcon		vcon
	velo		velo
	zone		zone

## 9.2 Code Execution

To run FEHM, the program executable file name is entered at the system prompt:

```
<PROMPT> xfehm
```

The I/O file information is provided to the code from an input control file or the terminal. The default control file name is *fehm.files*. If a control file with the default name is present in the directory from which the code is being executed, no terminal input is required. If the default control file is not present, input prompts are written to the screen. A short description of the I/O files used by FEHM precedes the initial prompt. The following assumes the default control file was not found in the execution directory (for this example /home/fehm/heat2d).

After the command **xfehm** is given, the code queries the user regarding the input files, as follows:

```
Enter name for iocntl -- default file name: not using
```

```
[(name/na or not using), RETURN = DEFAULT]
```

This query asks for a control file name. If a control file name is entered, no further terminal input is required. Figure 3 shows the control file that would produce the same results as the terminal responses discussed below and illustrated in Fig. 4.

```
/home/fehm/heat2d/input/heat2d.in
/home/fehm/heat2d/input/heat2d.in
/home/fehm/heat2d/input/heat2d.in
/home/fehm/heat2d/output/heat2d.out

/home/fehm/heat2d/output/heat2d.fin
/home/fehm/heat2d/output/heat2d.his

/home/fehm/heat2d/output/heat2d.chk
some
0
```

**Figure 3. Input control file for heat-conduction example.**

Files that are not needed for output can be represented with a blank line. If names are not provided for the write file or the data check file, the code will use the following defaults: *fehm.fin* and *fehm.chk*. Following the file names is the flag that controls terminal output. The last line of the file is the user subroutine number. Omitting these values results in no terminal output and no user subroutine call. For now, we assume a carriage return <cr> is entered and a control file is not being used. The following query will appear:

Enter name for inpt -- default file name: fehmn.dat

[(name/na or not using), RETURN = DEFAULT]

This query asks for an input file name. If a <cr> is given, the default *fehmn.dat* is used for the input file. We shall assume that the input file name entered is

input/heat2d.in

Note that a subdirectory containing the file is also given. If the file did not exist, the code would repeat the prompt for an input file. Next, the code would query to determine if the prefix of the input file name (the portion of the name preceding the final "." or first space) should be used for code-generated file names.

Do you want all file names of the form input/heat2d.\* ? [(y/n), RETURN = y]

\*\*\* Note: If "y" incoor and inzone will equal inpt \*\*\*

A <cr> will produce files with identical prefixes, including the subdirectory. If the response is negative, the code will query for the names of all required files.

Assume we enter "n".

Enter name for incoor -- default file name: input/heat2d.in

[(name/na or not using), RETURN = DEFAULT]

(See Fig. 4 for the remaining file name queries.)

Next a query for terminal output appears.

tty output -- show all reference nodes, selected reference nodes, or none:  
[(all/some/none), RETURN = none]

An "all" reply prints out the primary-node information to the terminal at every time step. A "some" reply prints a selected subset of the node information. A reply of "none" suppresses all tty output with the exception of error messages printed if code execution is terminated abnormally or when the maximum number of iterations are exceeded. Assume we enter "some".

The next query concerns the subroutine USER. This subroutine is used for special purposes and is not available to the general user.

user subroutine number (provided to subroutine USER before every time step):  
[RETURN = none]

Assume a <cr> is entered. The code will then print a summary of the I/O files to be used.

The final query regards the acceptance of the file set just created. A "yes" reply denotes that the user has accepted the file set and the code proceeds with calculations. A "no" reply starts the query sequence again so I/O file names may be reentered or modified. A "stop" reply stops the current computer job.

If data is OK enter yes to continue, no to restart terminal input,  
or stop to end program: [(yes/no/stop), RETURN = yes]

Screen output for this example execution using terminal input is shown in Fig. 4. User responses are shown in *italics*.

<PROMPT> *xfehm*

Version FEHMN XX-XX-XX 94/01/11 09:24:04

\*\*\*\* Default names for I/O files \*\*\*\*

control file	: fehm.files
input file	: filen.*
geometry data file	: filen.*
zone data file	: filen.*
output file	: filen.out
read file (if it exists)	: filen.ini
write file (if it exists)	: filen.fin
history plot file	: filen.his
tracer history plot file	: filen.trc
contour plot file	: filen.con
dual or dpdp contour plot file	: filen.dp
stiffness matrix data read/write file	: filen.stor
input check file	: filen.chk

\*\*\*\* where \*\*\*\*

"filen.\*" may be 100 characters maximum. If a name is not entered when prompted for, a default file name is used. "fehm.dat" is the default used for the input file name.

\*\*\*\* note \*\*\*\*

A save file and input check file are always written. If you do not provide a name for these files, the following defaults will be used: fehm.fin, fehm.chk

Enter name for iocntl -- default file name: not using

[(name/na or not using), RETURN = DEFAULT]

<cr>

Enter name for inpt -- default file name: fehm.dat

[(name/na or not using), RETURN = DEFAULT]

*input/heat2d.in*

Do you want all file names of the form example/heat2d.\* ? [(y/n), RETURN = y]

\*\*\* Note: If "y" incoor and inzone will equal inpt \*\*\*

*n*

Enter name for incoor -- default file name: input/heat2d.in

[(name/na or not using), RETURN = DEFAULT]

<cr>

**Figure 4. Terminal query for FEHM example run .**

Enter name for inzone -- default file name: input/heat2d.in

[(name/na or not using), RETURN = DEFAULT]  
<cr>

Enter name for iout -- default file name: input/heat2d.out

[(name/na or not using), RETURN = DEFAULT]  
*output/heat2d.out*

Enter name for iread -- default file name: input/heat2d.ini

[(name/na or not using), RETURN = DEFAULT]  
*na*

Enter name for isave -- default file name: input/heat2d.fin

[(name/na or not using), RETURN = DEFAULT]  
*output/heat2d.fin*

Enter name for ishis -- default file name: input/heat2d.his

[(name/na or not using), RETURN = DEFAULT]  
*output/heat2d.his*

Enter name for istrc -- default file name: input/heat2d.trc

[(name/na or not using), RETURN = DEFAULT]  
*na*

Enter name for iscon -- default file name: input/heat2d.con

[(name/na or not using), RETURN = DEFAULT]  
*na*

Enter name for iscon1 -- default file name: input/heat2d.dp

[(name/na or not using), RETURN = DEFAULT]  
*na*

Enter name for isstor -- default file name: input/heat2d.stor

[(name/na or not using), RETURN = DEFAULT]  
*na*

Enter name for ischk -- default file name: input/heat2d.chk

[(name/na or not using), RETURN = DEFAULT]  
*output/heat2d.chk*

**Figure 4. Terminal query for FEHM example run (continued).**

tty output -- show all reference nodes, selected reference nodes, or none:  
[(all/some/none), RETURN = none]  
*some*

user subroutine number (provided to subroutine USER before every time step):  
[RETURN = none]  
<cr>

First reference output node will be written to tty

File purpose - Variable - Unit number - File name

control	- iocntl	- 0	- not using
input	- inpt	- 11	- input/heat2d.in
geometry	- incoor	- 11	- input/heat2d.in
zone	- inzone	- 11	- input/heat2d.in
output	- iout	- 14	- output/heat2d.out
initial state	- iread	- 0	- not using
final state	- isave	- 16	- output/heat2d.fin
time history	- ishis	- 17	- output/heat2d.his
time his.(tr)	- istrc	- 18	- not using
contour plot	- iscon	- 19	- not using
con plot (dp)	- iscon1	- 20	- not using
fe coef stor	- isstor	- 21	- not using
input check	- ischk	- 22	- output/heat2d.chk

Value provided to subroutine user: not using

If data is OK enter yes to continue, no to restart terminal input,  
or stop to end program: [(yes/no/stop), RETURN = yes]  
<cr>

**Figure 4. Terminal query for FEHM example run (continued).**



### 9.3 Heat Conduction in a Square

This simple 2-D problem is used to illustrate input file construction and basic output. Heat conduction in a 1-meter square with an initial temperature,  $T_0 = 200^\circ\text{C}$ , is modeled after a surface temperature,  $T_s = 100^\circ\text{C}$ , is imposed at time,  $t = 0$  (Fig. 5). The input parameters used for the heat-conduction problem are defined in Table VIII. The finite-element mesh for this problem is shown in Fig. 6. Only a quarter of the square needs to be modeled because of problem symmetry.

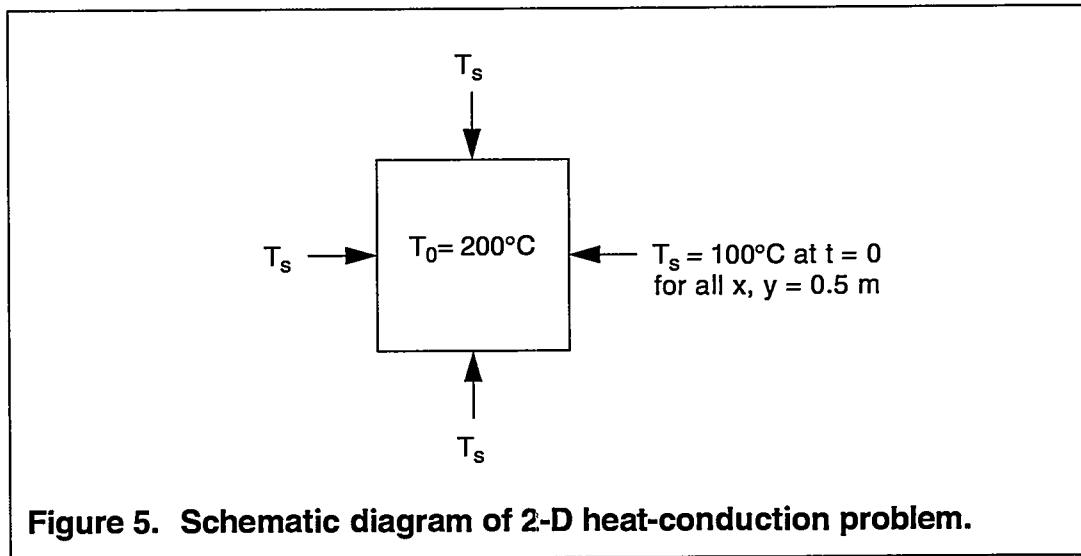
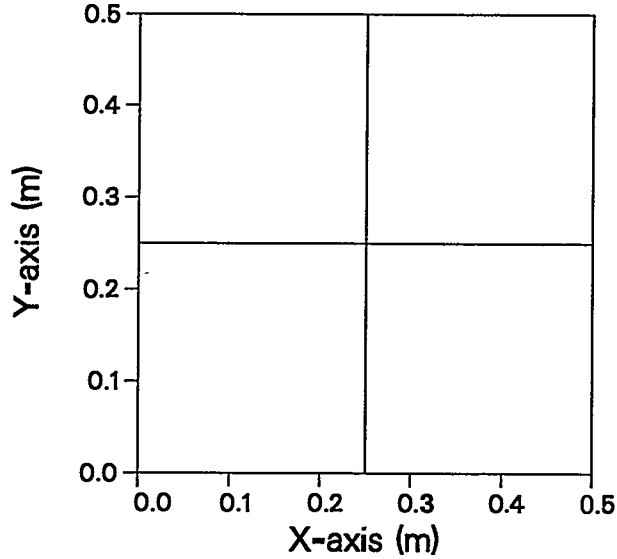


Table VIII. Input parameters for the 2-D heat-conduction problem		
Parameter	Symbol	Value
Rock thermal conductivity	$\kappa_r$	$2.7 \frac{\text{W}}{\text{m} \cdot \text{K}}$
Rock density	$\rho_r$	$2700 \text{ kg/m}^3$
Rock specific heat	$C_r$	$1000 \frac{\text{J}}{\text{kg} \cdot \text{K}}$
Width	$a$	0.5 m
Length	$b$	0.5 m
Initial temperature	$T_0$	$200^\circ\text{C}$
Surface temperature for all $x, y = 0.5 \text{ m}$	$T_s$	$100^\circ\text{C}$
Rock thermal diffusivity	$\kappa = \frac{\kappa_r}{\rho_r C_r}$	



**Figure 6. Finite-element mesh used for 2-D heat-conduction problem.**

The input file (see Fig. 7) uses the optional macro control statement **node** (output nodes) and the required macro control statements **sol** (solution specification - heat transfer only), **init** (initial value data), **rock** (rock properties), **cond** (thermal conductivities), **perm** (permeabilities), **time** (simulation timing data), **ctrl** (program control parameters), **coor** (node coordinates), **elem** (element node data), and **stop**. For this problem, the macro control statement **flow** is also used to set the temperature boundary conditions. A portion of the output file is reproduced in Fig. 8.

The analytical solution for 2-D heat conduction (Carslaw and Jaeger, 1959) is given by

$$T = T_s + \frac{16(T_0 - T_s)}{\pi^2} \sum_{m=0}^{\infty} \sum_{n=0}^{\infty} \frac{(-1)^{m+n}}{(2m+1)(2n+1)} \cos \frac{(2m+1)\pi x}{2a} \cos \frac{(2n+1)\pi y}{2b} e^{-\alpha_{m,n}t}$$

$$\text{where } \alpha_{m,n} = \frac{\kappa\pi^2}{4} \left[ \frac{(2m+1)^2}{a^2} + \frac{(2n+1)^2}{b^2} \right] \text{ and the region is taken to be}$$

$$-a < x < a, -b < y < b.$$

Figure 9 shows a plot of the simulation results compared to the analytical solution for the selected output nodes.

```

***** 2-D Heat Conduction Model (2X2 rectangles) *****
node
  2
  7 5
sol
  -1 -1
init
  10. 0. 200. 0. 0. 200. 0. 0.
rock
  1 9 1 2700. 1000. 0.

cond
  1 9 1 2.7e-00 2.7e-00 2.7e-00

perm
  1 9 1 1.e-30 1.e-30 1.e-30

flow
  1 3 1 10.00 -100.00 1.e03
  3 9 3 10.00 -100.00 1.e03

time
  0.005 4.00 1000 10 1994 02

ctrl
  40 1.e-04 08
  1 9 1 1

  1.0 0.0 1.0
  10 1.0 0.00005 0.005
  1 0
coor Feb 23, 1994 11:39:40
  9
  1 0. 0.50 0.
  2 0.25 0.50 0.
  3 0.50 0.50 0.
  4 0. 0.25 0.
  5 0.25 0.25 0.
  6 0.50 0.25 0.
  7 0. 0. 0.
  8 0.25 0. 0.
  9 0.50 0. 0.

elem
  4 4
  1 4 5 2 1
  2 5 6 3 2
  3 7 8 5 4
  4 8 9 6 5

stop

```

**Figure 7. FEHM input file for heat-conduction example (heat2d.in).**

FEHM 01.00 [sun4] 96/07/24 10:46:53

File purpose - Variable - Unit number - File name

control - iocntl - 0 - not using

input - inpt - 11 - heat2d.in

geometry - incoor - 11 - heat2d.in

zone - inzone - 11 - heat2d.in

output - iout - 14 - heat2d.out

initial state - iread - 0 - not using

final state - isave - 16 - heat2d.fin

time history - ish1s - 17 - heat2d.his

time his.(tr) - istrc - 0 - not using

contour plot - iscon - 0 - not using

con plot (dp) - iscon1 - 0 - not using

fe coef stor - isstor - 0 - not using

input check - ischk - 22 - heat2d.chk

Value provided to subroutine user: not using

\*\*\*\*\* 2-D Heat Conduction Model (2X2 rectangles) \*\*\*\*\*

\*\*\*\* input title : coor \*\*\*\* incoor = 11 \*\*\*\*

\*\*\*\* input title : elem \*\*\*\* incoor = 11 \*\*\*\*

\*\*\*\* input title : stop \*\*\*\* incoor = 11 \*\*\*\*

\*\*\*\* input title : node \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : sol \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : init \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : rock \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : cond \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : perm \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : flow \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : time \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : ctrl \*\*\*\* inpt = 11 \*\*\*\*

\*\*\*\* input title : stop \*\*\*\* inpt = 11 \*\*\*\*

BC to BC connection(s) found(now set=0.0)

**Figure 8. FEHM output from the 2-D heat-conduction example .**

```

pressures and temperatures set by gradients

storage needed for ncon      43 available      43
storage needed for nop       43 available      46
storage needed for a matrix   33 available      33
storage needed for b matrix   33 available      46
storage needed for gmres      81 available      81
storage available for b matrix resized to 33<<<<<<

time for reading input, forming coefficients 0.333E-01

**** analysis of input data on file fehmn.chk ****

*****
Time Step      1

      Timing Information
      Years      Days      Step Size (Days)
      0.136893E-04  0.500000E-02  0.500000E-02
Cpu Sec for Time Step = 0.1667E-01 Current Total = 0.1667E-01

      Equation Performance
Number of N-R Iterations:      1
Avg # of Linear Equation Solver Iterations: 3.0
Number of Active Nodes:      9.
Total Number of Newton-Raphson Iterations:      1
Node Equation 1 Residual Equation 2 Residual
  7  0.111444E-07      0.185894E-01
  5  0.165983E-07      0.135450E+01

      Nodal Information (Water)
                        source/sink source/sink
Node p(MPa) e(MJ) l sat temp(c) (kg/s) (MJ/s)
  7  10.000  0.00  0.000 199.981  0.  0.
  5  10.000  0.00  0.000 198.645  0.  0.

      Global Mass & Energy Balances
Total mass in system at this time:      0.000000E+00 kg
Total mass of steam in system at this time: 0.000000E+00 kg
Total enthalpy in system at this time:      0.105123E+03 MJ

Water discharge this time step: 0.000000E+00 kg (0.000000E+00 kg/s)
Water input this time step:      0.000000E+00 kg (0.000000E+00 kg/s)
Total water discharge:      0.000000E+00 kg (0.000000E+00 kg/s)
Total water input:      0.000000E+00 kg (0.000000E+00 kg/s)

Enthalpy discharge this time step: 0.297800E+02 MJ (0.689352E-01 MJ/s)
Enthalpy input this time step:      0.000000E+00 MJ (0.000000E+00 MJ/s)

```

**Figure 8. FEHM output from the 2-D heat-conduction example (continued).**

Total enthalpy discharge: 0.297800E+02 MJ (0.689352E-01 MJ/s)  
Total enthalpy input: 0.297800E+02 MJ (0.689352E-01 MJ/s)

Net kg water discharge (total out-total in): 0.000000E+00  
Net MJ discharge (total out-total in): 0.000000E+00  
Conservation Errors: 0.000000E+00 (mass), -0.100326E+01 (energy)

\*\*\*\*\*

Time Step 11

\*\*\*\*\*

Time Step 801

#### Timing Information

Years	Days	Step Size (Days)
0.109515E-01	0.400005E+01	0.500000E-04
Cpu Sec for Time Step = 0. Current Total = 4.533		

#### Equation Performance

Number of N-R Iterations: 1  
Avg # of Linear Equation Solver Iterations: 2.0  
Number of Active Nodes: 9.  
Total Number of Newton-Raphson Iterations: 801  
Node Equation 1 Residual Equation 2 Residual  
7 0.977369E-13 0.186062E-04  
5 0.621566E-13 0.930309E-05

#### Nodal Information (Water)

Node	p(MPa)	e(MJ)	Isat	temp(c)	source/sink (kg/s)	source/sink (MJ/s)
7	10.000	0.00	0.000	100.230	0.	0.
5	10.000	0.00	0.000	100.115	0.	0.

#### Global Mass & Energy Balances

Total mass in system at this time: 0.000000E+00 kg  
Total mass of steam in system at this time: 0.000000E+00 kg  
Total enthalpy in system at this time: 0.675565E+02 MJ

Water discharge this time step: 0.000000E+00 kg (0.000000E+00 kg/s)  
Water input this time step: 0.000000E+00 kg (0.000000E+00 kg/s)  
Total water discharge: 0.000000E+00 kg (0.000000E+00 kg/s)  
Total water input: 0.000000E+00 kg (0.000000E+00 kg/s)

Enthalpy discharge this time step: 0.455636E-05 MJ (0.105471E-05 MJ/s)  
Enthalpy input this time step: 0.000000E+00 MJ (0.000000E+00 MJ/s)  
Total enthalpy discharge: 0.673463E+02 MJ (0.155894E+02 MJ/s)

**Figure 8. FEHM output from the 2-D heat-conduction example (continued).**

Total enthalpy input: 0.673463E+02 MJ (0.155894E+02 MJ/s)  
Net kg water discharge (total out-total in): 0.000000E+00  
Net MJ discharge (total out-total in): 0.000000E+00  
Conservation Errors: 0.000000E+00 (mass), -0.100144E+01 (energy)

simulation ended: days 4.00 timesteps 801  
total newton-raphson iterations = 801

total code time(timesteps) = 4.483334

```

****-----****
**** This program for ****
**** Finite Element Heat and Mass Transfer in porous media ****
****-----****
**** Version : FEHM 01.00 [sun4] ****
**** End Date : 96/07/24 ****
**** Time : 10:46:59 ****
****-----****

```

Figure 8. FEHM output from the 2-D heat-conduction example (continued).

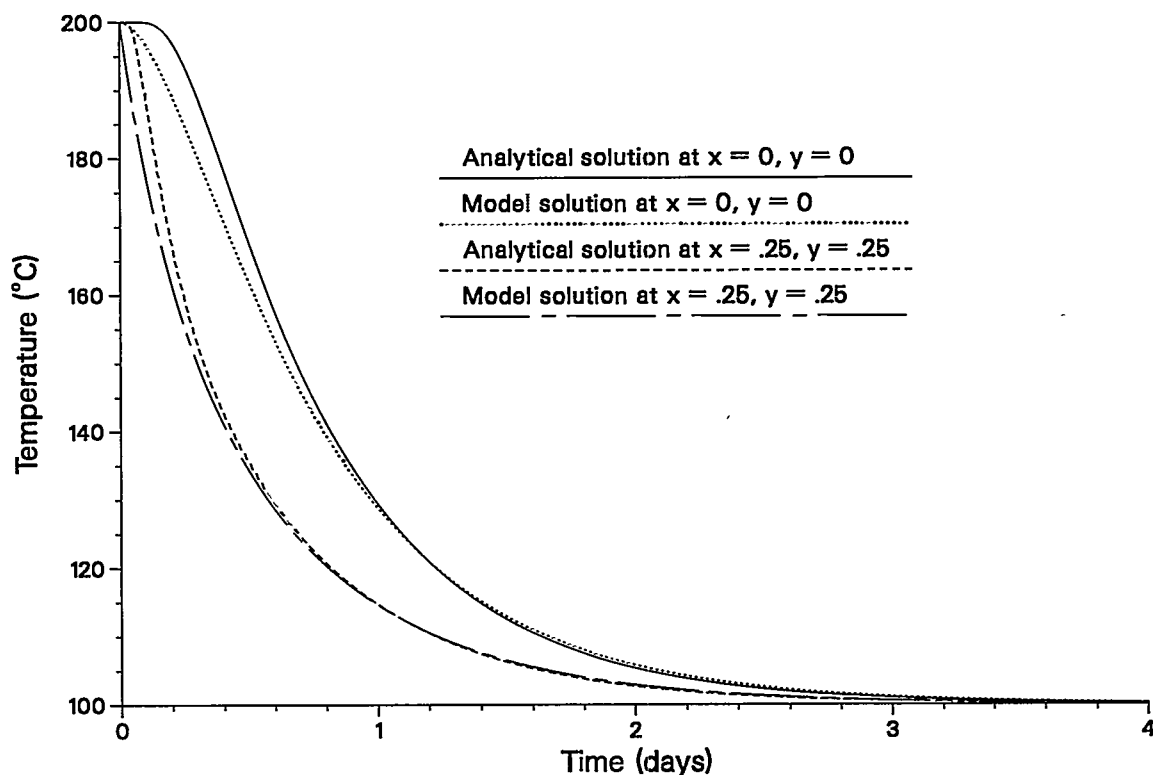


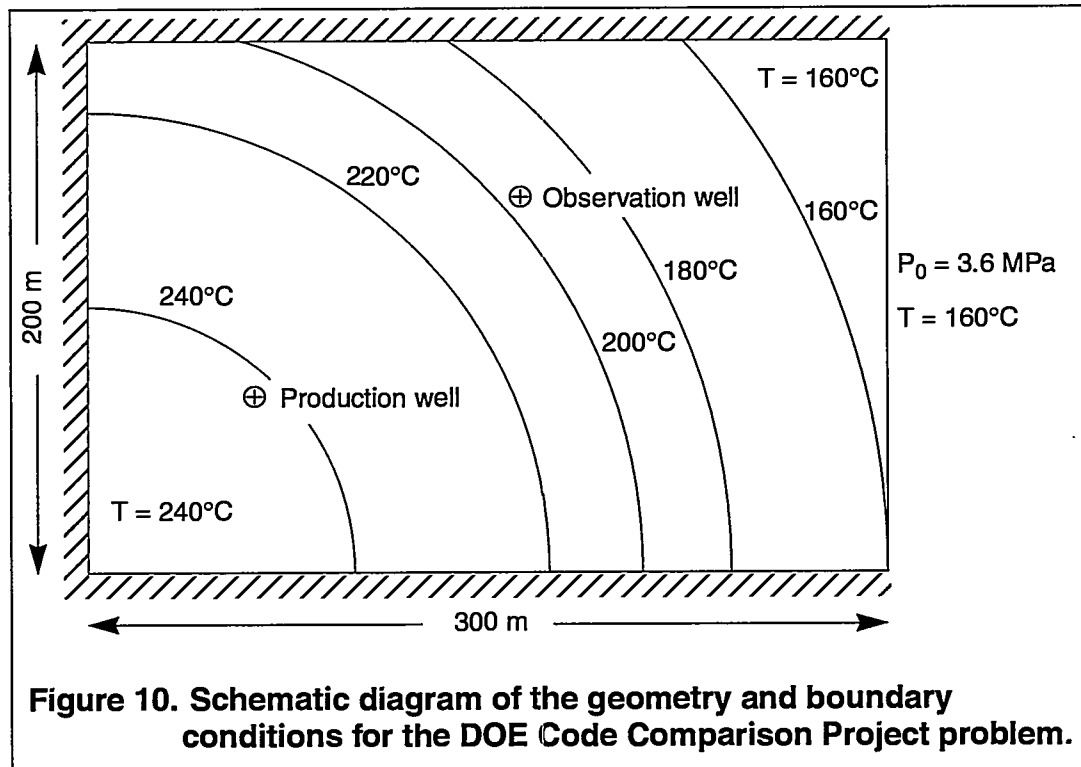
Figure 9. Comparison of analytical and model solution for 2-D heat conduction.

## 9.4 DOE Code Comparison Project, Problem 5, Case A

This problem involves multiphase flow in a 2-D horizontal reservoir. The problem is characterized by a moving two-phase region, i.e., the fluid produced at the production well is replaced by cold water recharge over one of the outer boundaries. The problem parameters are given in Table IX, and the geometry and boundary conditions are shown in Fig. 10. Of particular note are the variable initial-temperature field provided to the code through a read file (see Section 5.7 on page 10), and the prescribed pressure and temperature on the right boundary. A partial listing of the input file is provided in Fig. 11. In addition to the required macros, the macro **flow** is used to specify the pressure and temperature boundary condition and the production flow rate. Macro **rlp** is used to set the residual liquid and gas saturations.

Table IX. Input parameters for the DOE Code Comparison Project problem		
Parameter	Symbol	Value
Reservoir permeability	$k$	$2.5 \times 10^{-14} \text{ m}^2$
Reservoir porosity	$\phi$	0.35
Rock thermal conductivity	$\kappa_r$	$1 \frac{\text{W}}{\text{m} \cdot \text{K}}$
Rock density	$\rho_r$	$2563 \text{ kg/m}^3$
Rock specific heat	$C_r$	$1010 \frac{\text{J}}{\text{kg} \cdot \text{K}}$
Reservoir length	$x$	300 m
Reservoir thickness	$y$	200 m
Liquid residual saturation	$s_{lr}$	0.3
Gas residual saturation	$s_{gr}$	0.1
Reservoir discharge	$q_m$	$0.05 \frac{\text{kg}}{\text{m} \cdot \text{s}}$
Initial Pressure	$P_o$	3.6 MPa
Production well coordinates:	$x = 62.5 \text{ m}, y = 62.5 \text{ m}$	
Observation well coordinates:	$x = 162.5 \text{ m}, y = 137.5 \text{ m}$	
Initial temperature distribution ( $T$ in $^{\circ}\text{C}$ , $r$ in m):		
$T(x, y, 0) = \begin{cases} 240 & 0 \leq r \\ 240 - 160\left(\frac{r-100}{200}\right)^2 + 80\left(\frac{r-100}{200}\right)^4 & 100 < r < 300 \\ 160 & r \geq 300 \end{cases}$		
where $r = \sqrt{x^2 + y^2}$		





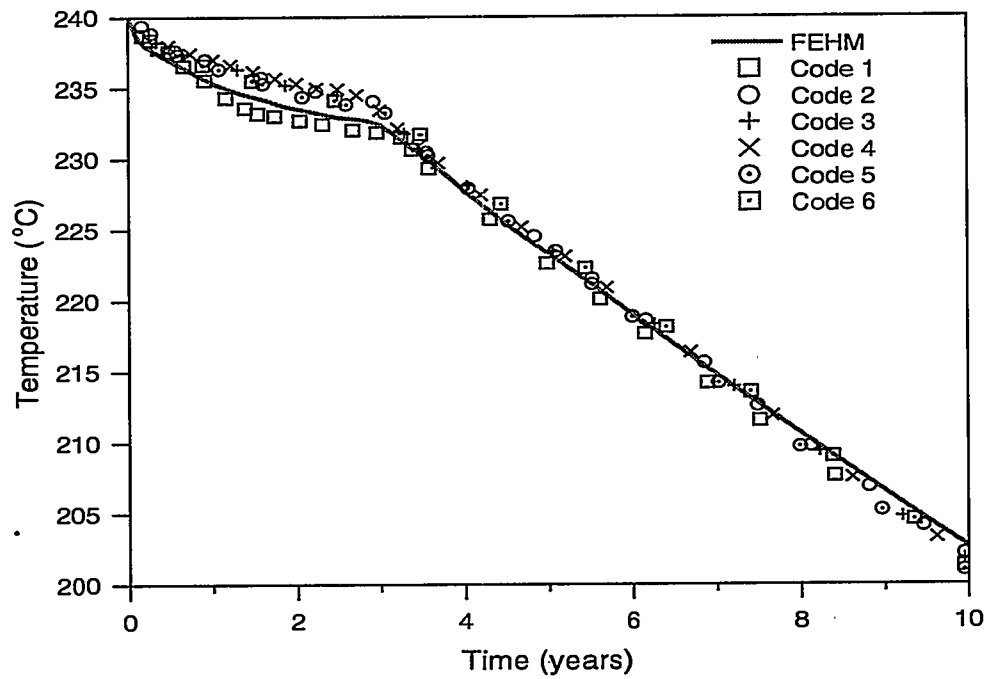
There is no analytical solution for this problem, but six researchers produced results for the DOE Code Comparison Project (Molloy 1980). The reader is referred to this reference for a more detailed discussion of this problem and the code comparison. Results from this problem are compared to those for the other codes, obtained from Molloy (1980), as a check on FEHM. The results for the outlet temperature, shown in Fig. 12, are in excellent agreement with the other codes. The results for the outlet pressure and pressure at an observation well 125 m distant (Fig. 13) are also in good agreement with the other codes. Contour plots of pressure and temperature at the end of the simulation were also generated for this problem and are shown in Fig. 14 and Fig. 15.

```

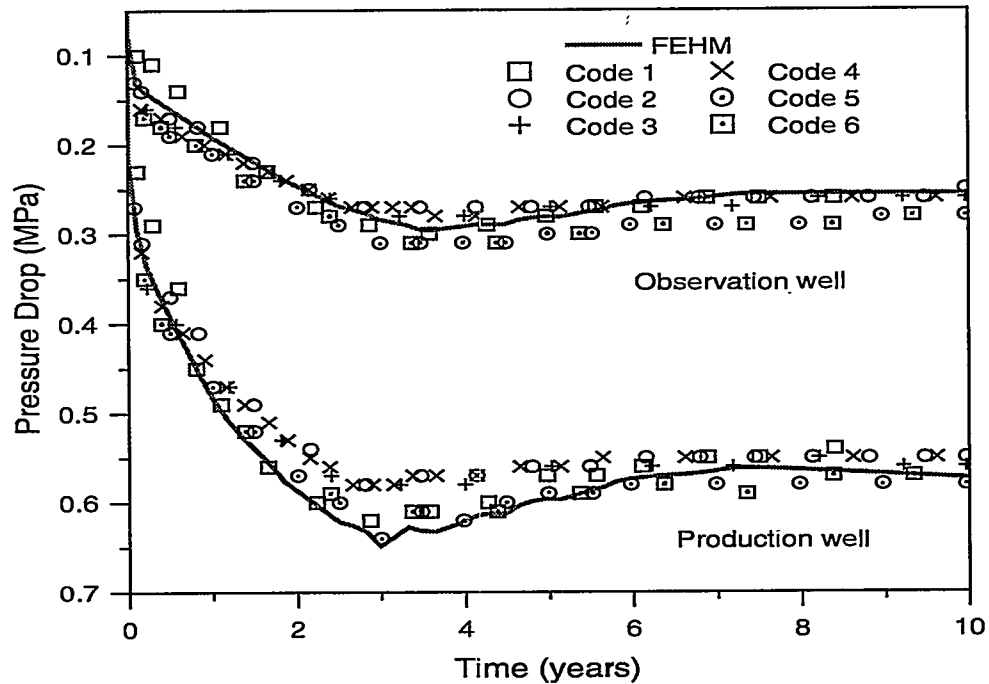
*** DOE Code Comparison Project, Problem 5, Case A ***
node
  2
  50 88
sol
  1 1
init
  3.6 0.240. 0.0. 240.0. 0.
rlp
  2 0.30.1 0.00.0
  1 140 1 1
rock
  1 140 1 2563. 1010. 0.35
cond
  1 140 1 1.00e-00 1.00e-00 1.00e-00
perm
  1 140 1 2.5e-14 2.5e-14 0.e-00
flow
  88 88 1 0.050 -25.00 0.
  14 140 14 3.600 -160.00 1.
time
  30.03650.100001000199403
ctrl
  40 1.e-0708
  1 140 1 1
  1.0 0.01.0
  40 1.20.1 60.
  1 0
coord
  140
  .
  .
  .
elem
  4 117
  .
  .
  .
stop

```

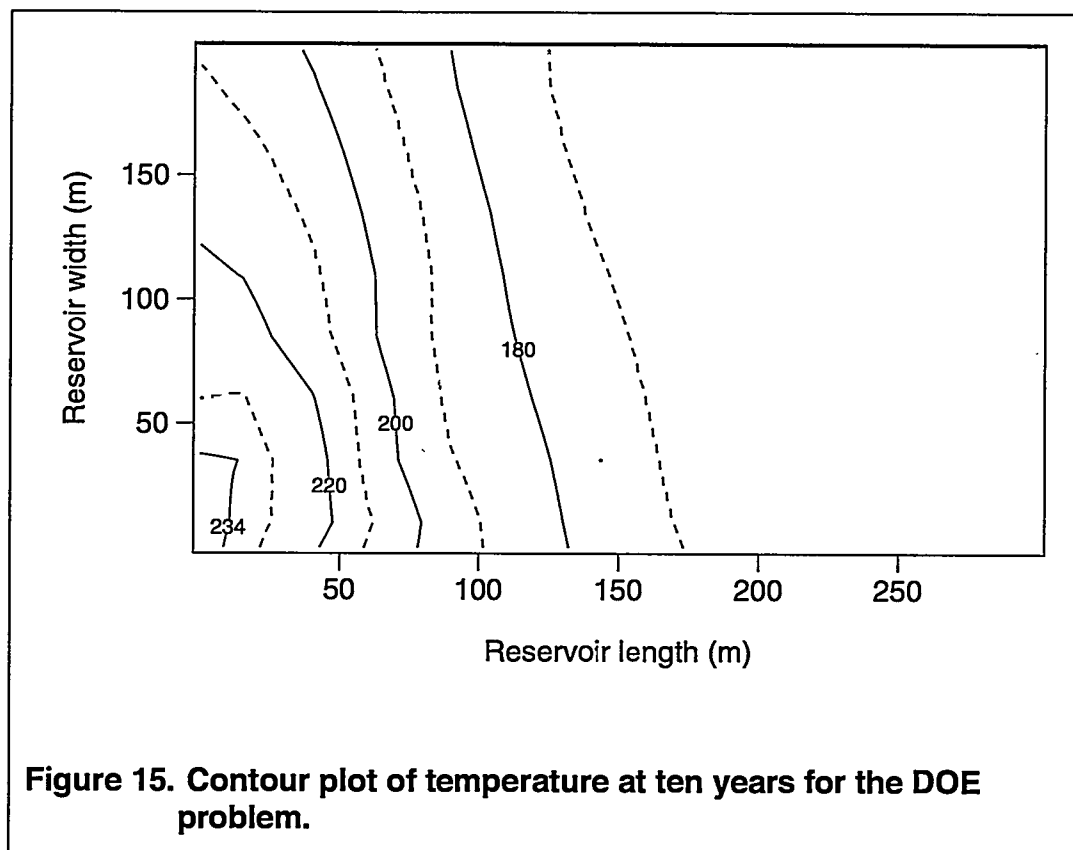
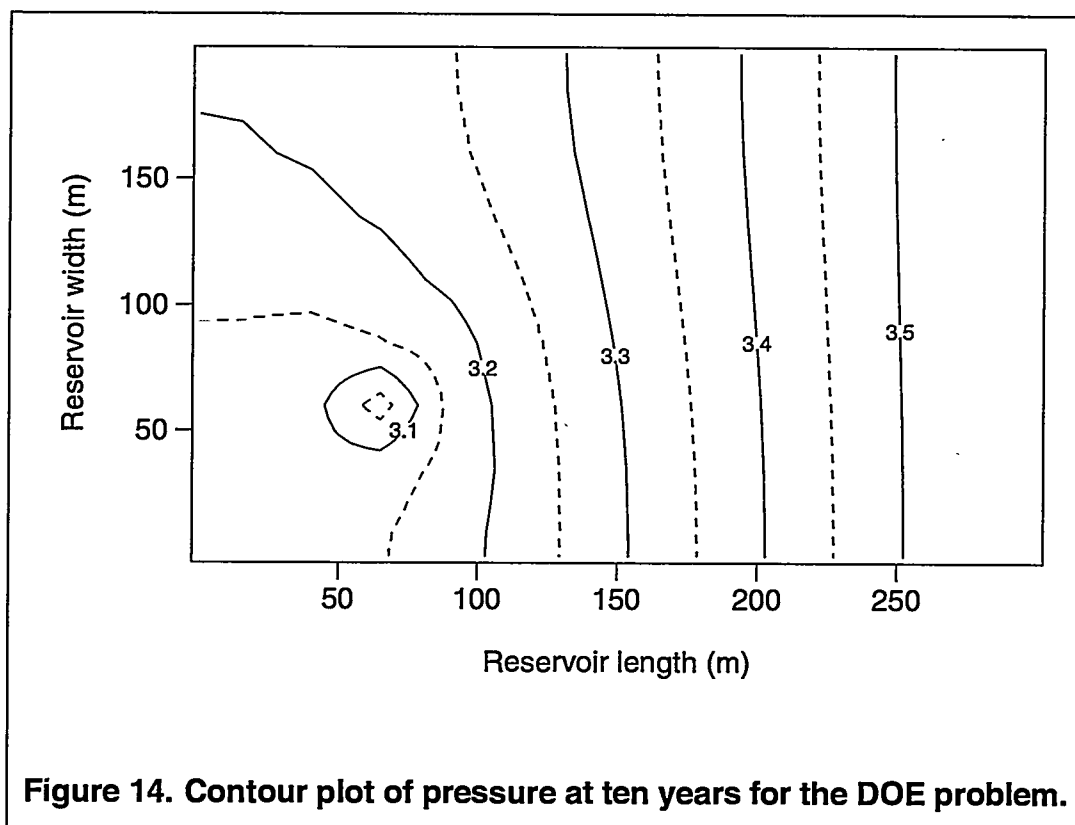
**Figure 11. FEHM input file for DOE problem.**



**Figure 12. Comparison of FEHM production-well temperatures with results from other codes.**



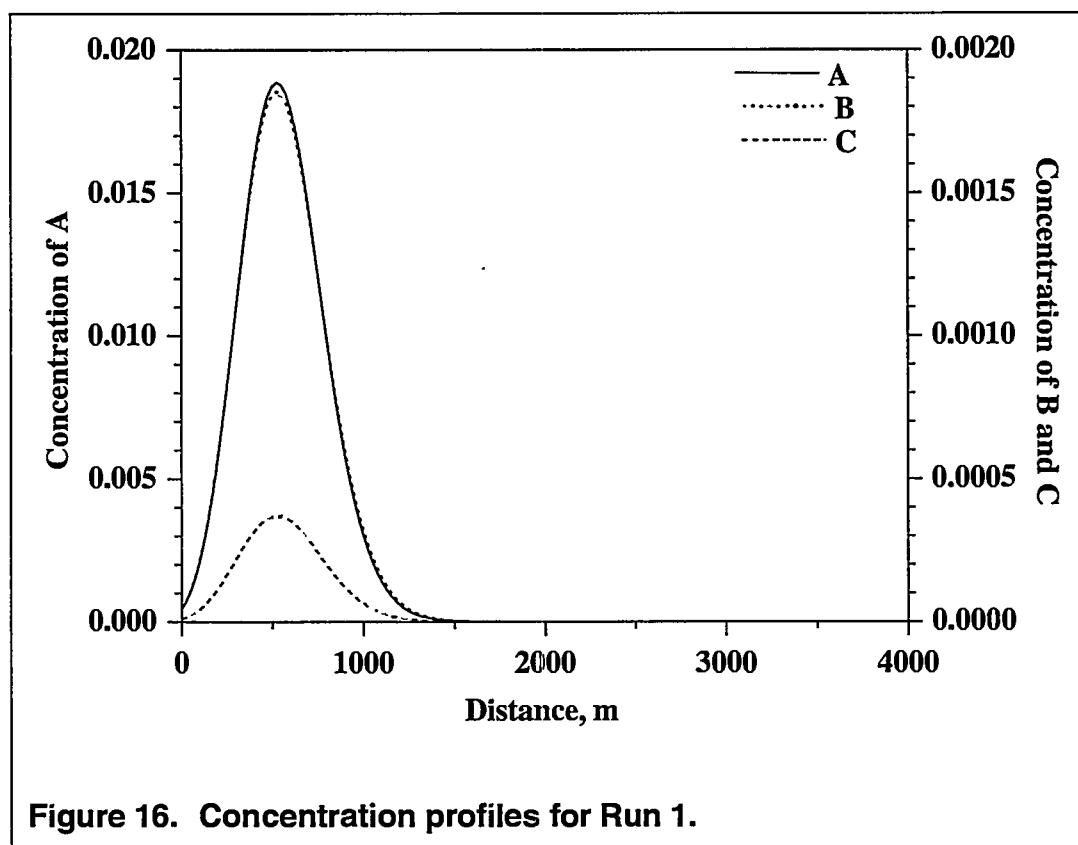
**Figure 13. Comparison of FEHM production- and observation-well pressure drops with results from other codes.**



## 9.5 Reactive-Transport Example

This one-dimensional example demonstrates the use of the reactive-transport module of FEHM. The input for the **rxn** and **trac** macros are those discussed in the example input for these macros in Section 6.2.44 and Section 6.2.54. The flow system is a one-dimensional flow path of 402 nodes ( $201 \times 2$ ) with rock properties and flow rates such that the mean fluid residence time in the path is 10,000 yr. Species A sorbs with a  $K_d$  of 0.1, which, for the rock properties chosen, is equivalent to a retardation factor of 5. The solute transport problem is run for 5,000 yr, or half of the mean residence time of the fluid. Therefore, in the absence of other reactions, species A would be expected to travel  $0.5/5 = 0.1$  of the length of the column.

When chemical reactions are included, the situation becomes more complex. Fig. 16 shows the concentration profiles at the end of the simulation for this example



(called Run 1). Even though the first reaction is specified as kinetically controlled, the rate constants are large enough for the reaction to virtually reach equilibrium over the time period of the simulation ( $k_r\tau = 50$ , where  $\tau$  is the time of the simulation). Thus, the concentrations of A and B essentially reach equilibrium; the equilibrium constant is the ratio of the rate constants, or 0.1 (there is 10 times as much A as B in solution). Of course, A is also present on the rock surface wherever concentrations are nonzero. Solute C travels with A and B and is in equilibrium

with *B* in solution; its concentration is 0.2 times that of *B* everywhere because of the equilibrium constant chosen. The entire suite of solutes has moved roughly 10% of the way down the column, as discussed above. With chemical reactions, the rate of movement of the solutes can be faster than that of a nonreacting species that only sorbs, because solutes *B* and *C* do not sorb. However, for this example, solute *A* constitutes about 90% of the aqueous portion of the contaminant. Kinetics and equilibrium parameters that favor the formation of *B* and *C* would in turn increase the rate of movement of the contaminants.

A common application is radionuclide migration, for which we must include radioactive decay of each solute. It may also be of interest to track the movement of the daughter product. In Run 2, we assume that all species of Run 1 are species of the same radionuclide, each of which undergoes decay to a long-lived, mobile radionuclide that does not sorb. The input file is changed to include a fourth solute in the **trac** macro that is identical in input to the second and third solutes. The changes to the **rxn** macro are more complex and thus are shown in Fig. 17.

rxn							
group							
2							
1	1	1	0				
0	0	0	1				
5	0						
kinetic	3.1688e-11	0.	3.1688e-10	0.			
equilibrium	1	0.2	0.	1.e-2	1.e-3	1.e-10	
kinetic	3.1688e-12	0.	0.	0.			
kinetic	3.1688e-12	0.	0.	0.			
kinetic	3.1688e-12	0.	0.	0.			
1	-1	0	0				
0	1	-1	0				
1	0	0	-1				
0	1	0	-1				
0	0	1	-1				
1	-1	0	0				
0	1	-1	0				
1	0	0	-1				
0	1	0	-1				
0	0	1	-1				
1	1	1	1				
1	1	1	1				
1	1	1	1				
1	1	1	1				
1	1	1	1				
0	0	0	0				
0	0	0	0				
1	1	1	1				
0	0	0	0				
0	0	0	0				

Figure 17. FEHM macro rxn in the input file for reactive-transport example.

Radioactive decay is handled by adding three reactions, one each for *A*, *B*, and *C* reacting to form *D*. The kinetics of each of these reactions are identical, reflecting an irreversible, first-order decay to form the daughter product. Solute *A* sorbs to the rock surface; therefore, we must specify that the decay reaction takes place for both the aqueous and solid portions of the solute. This is done by setting *FL\_MULT* and *SB\_MULT* to 1 for the third reaction.

Figure 18 shows the results of this example. A significant amount of daughter

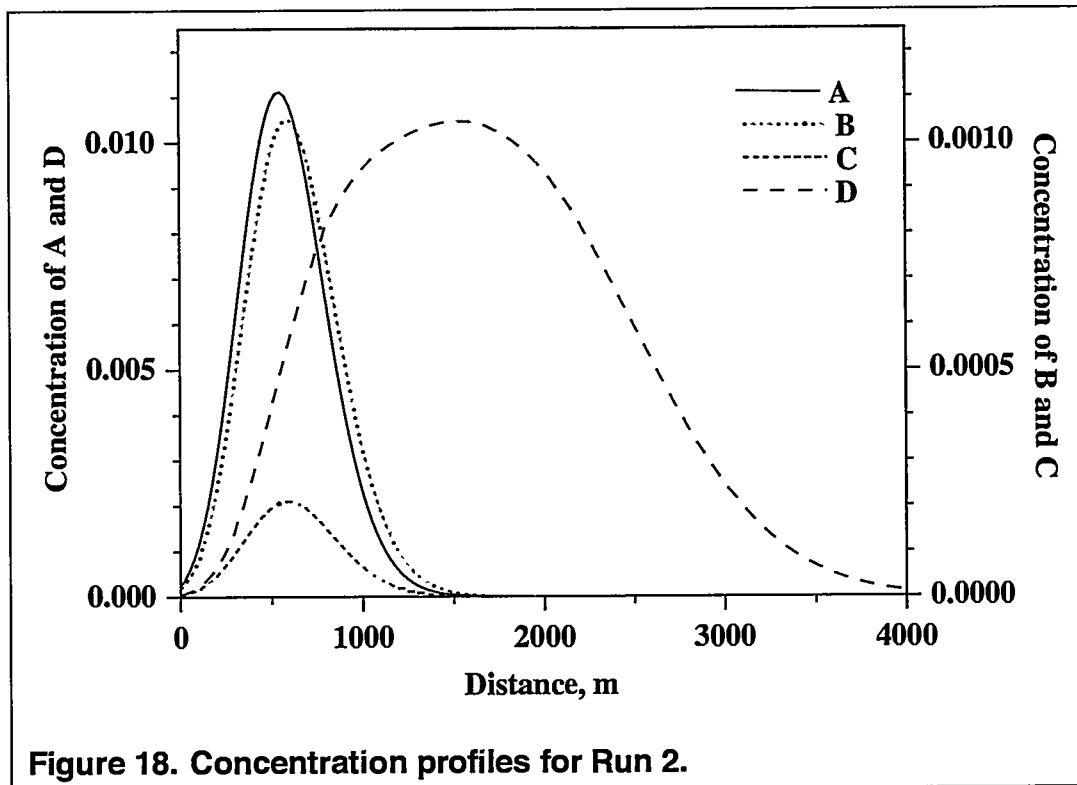


Figure 18. Concentration profiles for Run 2.

product *D* has been produced, and it is not forced to travel at the reduced velocity of sorbing solute *A* because it is connected to the other solutes only through the radioactive decay source term. The original solutes behave as they did in Run 1, except that their concentrations are somewhat reduced due to radioactive decay.

**Considerations of Numerical Efficiency.** The decision of how to group the solutes represents a trade-off among the robustness required for a given reactive transport system, memory requirements, and computational speed. We use these example runs to illustrate some of the considerations. In the discussions below, when we speak of the coupling of solutes, we refer to the method of grouping the solutes into systems of equations that are solved simultaneously. Regardless of the grouping of and order in which the solute concentrations are solved, the code requires that the full system of interacting solutes reach convergence at every time step. The overall solution is therefore “fully coupled,” regardless of the details of the solution procedure.

In Run 1, we solved first for solute *A* alone, after which solutes *B* and *C* were coupled. Solutes that are coupled only through a kinetic reaction need not be solved simultaneously as long as the kinetics are not too rapid. When systems are solved as more than one group, the time required to complete one outer iteration is shorter, but more outer iterations will be required (only one outer iteration is required if all solutes are coupled into a single group). A rule of thumb is that as long as only a few outer iterations are required, solving the problem in several groups will be competitive with a more fully coupled solution. For example, in Run 1, two or three outer iterations were typically required. Coupling all three solutes reduces the number of outer iterations to 1, but the computational time was virtually identical in this example problem. Note that solutes coupled through equilibrium reactions must always be solved simultaneously because these reactions are specified, within the code, with very rapid kinetics to approximate equilibrium behavior. An added benefit of solving the system as several groups is that the memory requirements are lower. This factor was not a consideration for this example, which solved a system of only 402 nodes.

In Run 2, notice that the fourth solute was solved alone subsequent to a group that coupled the first three. There is no benefit to the convergence of the system of equations from simultaneously solving solute *D* with the other three because it is formed only from irreversible reactions (radioactive decay) involving solutes *A*, *B*, and *C*. This fact means that there is no "feedback" from the concentration of *D* onto the other concentrations. The decay reactions provide the source term for solute *D*, but the concentration of *D* does not impact the solution of the other solutes; the coupling is one-way. The only potential benefit is that when all solutes are coupled, the system is automatically solved in a single outer iteration, whereas the code cannot assume overall convergence and must perform a second outer iteration when *D* is decoupled from the other three solutes. For this problem, the benefit of fully coupling the solution is almost exactly counterbalanced by the additional work of finding a four degree-of-freedom solution (versus three followed by one), so that the two solutions take comparable times to finish.



## 10.0 USER SUPPORT

Licensing and installation support can be received from:

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## APPENDIX: FEHM VERIFICATION SCRIPTS

### A. DESCRIPTION OF SCRIPTS

A series of *cshell* scripts were developed to perform the verification operations (see Table X). A primary script, **FEHM V&V Script for Execution of Comparison Tests (FEHM\_VVSECT)**, controls the FEHM verification runs via an execution script (**RUN\_VERIFICATION**) then generates a comparison of results and summary via a supporting comparison tests script (**COMPARE\_RESULTS**). An execution log is generated when the primary script is run (see Sections E through G for listings of the three top-level scripts and Sections H and I for examples of the specific problem execution and run comparison scripts). In addition to being invoked by the **FEHM\_VVSECT** script, the **RUN\_VERIFICATION** and **COMPARE\_RESULTS** scripts may be invoked independently to re-execute specific tests or comparisons by providing problem descriptors as arguments. Two subsidiary programs, **COMPARE** and **SUMMARIZE**, used by the comparison tests script, are also described in Table X.

Table X. Scripts and support programs for verification operations	
Script name	Description
FEHM_VVSECT	Executes all verification test problems ( <b>RUN_VERIFICATION</b> ) and generates comparison of results and summary ( <b>COMPARE_RESULTS</b> ) usage: <b>FEHM_VVSECT</b> [xfehmn_path verification_dir]
RUN_VERIFICATION	Invokes script for each verification test problem ( <b>runproblem</b> ) usage: <b>RUN_VERIFICATION</b> [-p xfehmn_path verification_dir] [problem1 problem2 ...]
COMPARE_RESULTS	Invokes script to generate result comparison and summary for each verification test problem ( <b>compproblem</b> ) usage: <b>COMPARE_RESULTS</b> [-p verification_dir] [-i date_id] [problem1 problem2 ...]
runproblem	Problem-specific script for <b>FEHM</b> input/output setup and code execution
compproblem	Problem-specific script for <b>COMPARE</b> and <b>SUMMARIZE</b> input/output setup and execution
MAKE_OUTPUT_DIRS	Script for making output directories for verification problems in a user's local verification directory
COMPARE	Program that reads <b>FEHM</b> results and generates a numeric comparison with analytical or alternate code solutions.
SUMMARIZE	Program that reads results from <b>COMPARE</b> and outputs the results for related groups of tests in a single table.

The problems are set up in a directory-tree file structure. The root/verification directory contains the primary (**FEHM\_VVSECT**), execution (**RUN\_VERIFICATION**), and comparison (**COMPARE\_RESULTS**) scripts with a subdirectory for each test problem.

The execution log and summary report are written to the root directory (see Sections J and K). Each problem directory contains the problem-specific execution (**runproblem**) and comparison (**compproblem**) scripts, an input and output subdirectory, and other files and directories as needed (Fig. 19). The following problems are currently run by the **FEHM\_VVSECT** script (the section numbers in parentheses correspond to the problem and result descriptions found in Dash et al. 1997): avdonin (5.8), dissolution (5.15), doe5a (5.10), dryout (5.11), dual (5.7), fracture\_transport (5.14), heat2d (5.2), heat3d (5.2), henrys\_law (5.13), infiltration (5.5), multi\_solute (5.16), ramey (5.3), sorption (5.12), theis (5.4), toronyi (5.9), transport3D (5.17), and vapor\_extraction (5.6). As additional test problems are developed, they will be incorporated into the test environment. The thermodynamics tests (5.1) are run independently because the functions need only be retested if the polynomial coefficients are modified. Also, any errors introduced to the routines containing the thermodynamic functions would result in errors in the other tests.

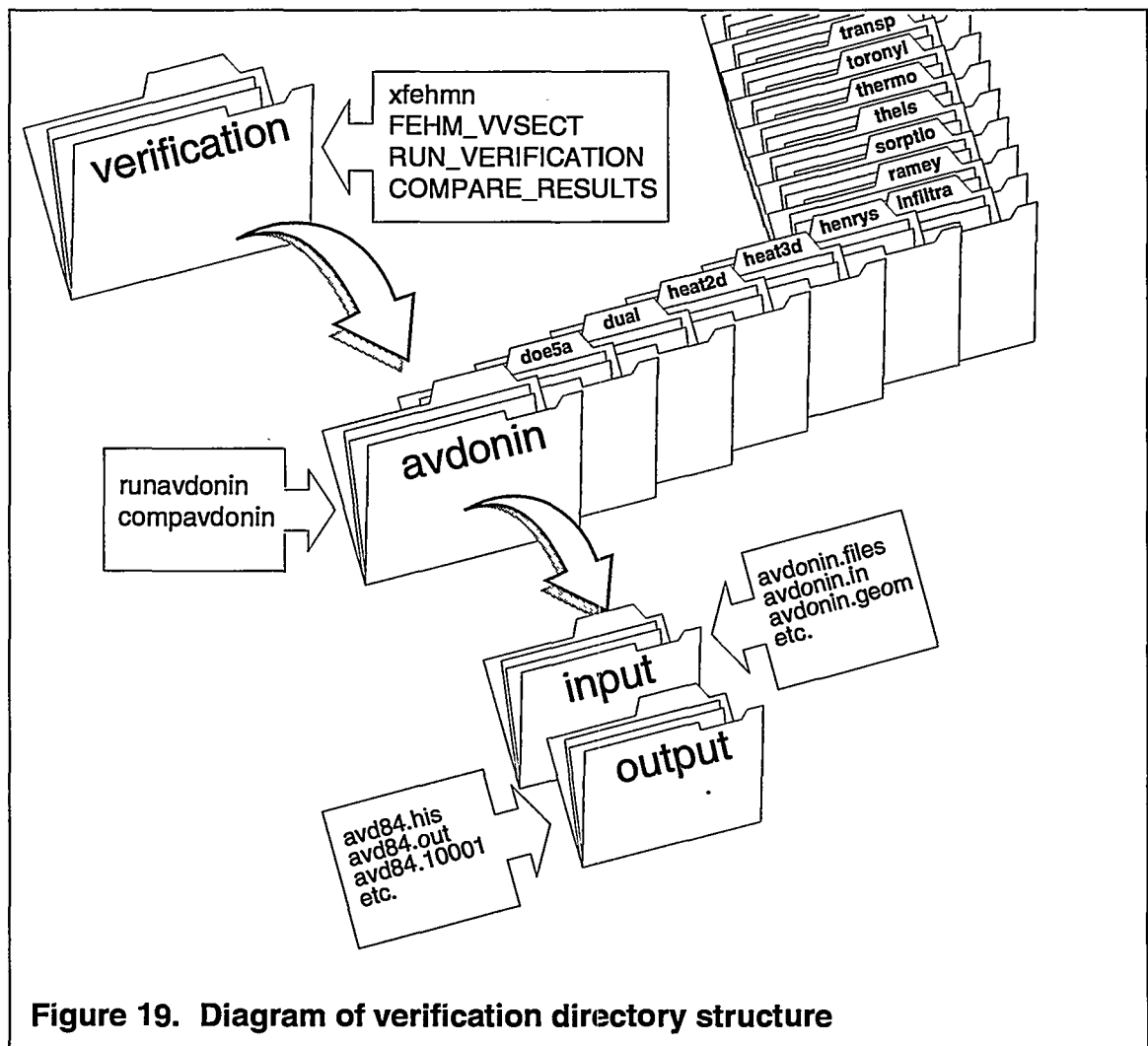


Figure 19. Diagram of verification directory structure

## B. INSTALLATION

Files needed to set up the verification environment are contained in a tar file called `verification.tar`. A directory should be created and the files installed there, i.e.,

```
% mkdir verification
% cd verification
% tar xvf directory/verification.tar
```

where *directory* is the location of the `verification.tar` file.

When the files have been installed in the verification directory, the auxiliary programs need to be built and installed. To do this, change into verification subdirectory `SRC`. Currently, there are two subdirectories under `SRC`: `compare` and `thermodynamics`. The `compare` subdirectory contains programs (`COMPARE`, `COMPARET`, and `SUMMARIZE`) that are used to compare FEHM results to analytical and other model solutions and to summarize the comparisons. The `thermodynamics` subdirectory contains programs (`COMPSAT` and `COMPETHER`) that use the FEHM thermodynamic functions to generate values over the valid range of use. The programs require that the `SRCDIR` environment variable in each Makefile be set to the directory that contains the FEHM source code so that source for the malloc and parser routines may be found. In addition, program `COMPSAT` uses source code for the `psat` function and common files `comai.h` and `comdti.h`.

To build programs `COMPARE`, `COMPARET`, and `SUMMARIZE`:

```
% cd SRC/compare
% make -f Makefile.compare machine
```

where *machine* is `sun`, `hp`, `ibm`, `sgi`, or `cray`.

Links from the `thermodynamics` subdirectory may be made to `comai.h` and `comdti.h` of the FEHM source code to use the most up-to-date versions of these files during compilation or the files may be copied to the `thermodynamics` subdirectory. Common file `comii.h` has been modified for use with these test programs so the version in the FEHM objects directory currently cannot be used. To build programs `COMPSAT` and `COMPETHER`:

```
% cd SRC/thermodynamics
% make -f Makefile.thermo all
```

The verification scripts expect the auxiliary program binaries to reside at the top level of the verification directory. This requirement can be accomplished either by moving the binaries there or by making links to their actual location if they are installed elsewhere.

## C. USING THE VERIFICATION SCRIPTS

To run the verification tests the user should change to the directory where the output should reside. (For the owner of the verification directory, this is usually the verification directory.) This directory should contain a subdirectory for each verification

problem to be run and each problem subdirectory should contain an output subdirectory. A script called `MAKE_OUTPUT_DIRS`, located in the verification directory, can be used to create (in the current directory) a directory and output subdirectory for each problem. The user has two options for defining the location of the verification directory (the directory containing verification scripts, problem input files, etc.) and the FEHM executable. The variables may either be entered as command line arguments to the test scripts (see "usage" in Table X) or they may be defined in a file called `PATHS` in the directory from which the scripts will be executed (see Fig. 20). Variables defined on the

```
#!/bin/csh
#   PATHS

# Set location where FEHM executable can be found:

setenv XFEHMN /home/fehm/bin/xfehm.V01.00

# Set location of verification directory.
# Verification scripts and executables should be found here, along with
# problem subdirectories and their associated scripts and input files, i.e.,
# VER_DIR/problem_dir/input, output directories may be located here
# or elsewhere

setenv VER_DIR /home/fehm/verification
```

**Figure 20. Example of PATHS file in script-execution directory**

command line have precedence over values set in the `PATHS` file. Complete path names must be used.

To run all verification test problems use **FEHM\_VVSECT**, i.e.,

```
% FEHM_VVSECT (assumes PATHS file is present)
```

or

```
% FEHM_VVSECT -p ~/bin/myxfehmn /home/fehm/verification
```

The execution directory will contain a log of the runs and result comparisons in files called `VERIFICATION.date` and `SUMMARY_RPT.date`, where *date* is the date of execution in yymmdd format (see Sections J and K for an example of these files). It should be noted that if a log file or summary file with the current date or identifier already exists in the execution directory it will be renamed, i.e., `VERIFICATION.date.old` or `SUMMARY_RPT.date.old`.

To run selected problems use **RUN\_VERIFICATION** and **COMPARE\_RESULTS**, i.e.,

```
% RUN_VERIFICATION thermodynamics
```

```
% COMPARE_RESULTS -i thermodynamics thermodynamics
```

or

```
% RUN_VERIFICATION -p ~/bin/myxfehmn /home/fehm/verification doe5a dual
```

`% COMPARE_RESULTS doe5a dual`

For the first case, the summary is found in `SUMMARY_RPT.thermodynamics`, whereas for the second case, it is found in `SUMMARY_RPT.date` because an identifier was not specified. The execution sequence is logged to the terminal unless output is redirected when the scripts are invoked.

If `RUN_VERIFICATION` and `COMPARE_RESULTS` are invoked without arguments, all test problems and comparisons will be run.

## **D. ASSUMPTIONS AND LIMITATIONS**

The verification scripts were developed on a Sun-4 architecture and have been tested on HP, IBM, SGI, and Sun, but they should work on any standard UNIX workstation. The examples provided below were run on a Sun.

## E. FEHM\_VVSECT

### FEHM\_VVSECT script

```
#!/bin/csh
# FEHM_VVSECT
# FEHM V&V Script for Execution of Comparison Tests

set date=`date +%y%m%d`
set results=VERIFICATION.$date

if (-f $results) then
    mv $results $results.old
endif

@ paths = 0
# If the PATHS are present as command line arguments
if ($#argv == 2) then
    setenv XFEHMN $1
    setenv VER_DIR $2
    @ paths = 1

# or If the PATHS file exists set the executable and directory paths
else if (-f PATHS) then
    source PATHS
    @ paths = 2

else
    echo 'PATHS to verification executables must be set.'
    echo 'They can be entered as command line arguments or put in a file'
    echo 'called PATHS in the current directory.'
    echo ' '
    echo 'usage: FEHM_VVSECT [xfehmn_path verification_dir]'
    exit 1
endif

@ flag = 0
# Verify that XFEHMN is executable
if (! -x $XFEHMN) then
    echo '$XFEHMN' does not exist or is not an executable file.'
    @ flag = 1
endif
# Verify that VER_DIR is a directory
if (! -d $VER_DIR) then
    echo '$VER_DIR' is not a directory file.'
    @ flag = 1
endif
```

### FEHM\_VVSECT script (continued)

```
if ($flag != 0) then
    if ($paths == 2) then
        echo 'Check your PATHS file'
    else
        echo 'Check your command line input'
    endif
    exit 1
endif

# Execute FEHMN with no input to determine version being tested
touch fehmn.files
$XFEHMN >& /dev/null; wait
set version=`cat fehmn.chk | nawk '{print $1" "$2}'`
rm fehmn.*

echo $version: Verification started "`date`" >& $results
echo ' ' >>& $results

# Execute FEHMN verification problems
if (-e $VER_DIR/RUN_VERIFICATION) then
    RUN_VERIFICATION >>& $results &; wait
    @ return = $status
    if ($return != 0) then
        echo 'Status: '$return' RUN_VERIFICATION exited with ERROR'
    endif
    echo ' ' >>& $results
else
    echo "Can't find RUN_VERIFICATION - check your verification directory"
    exit 1
endif

# Run problem comparisons and summarize results
if (-e $VER_DIR/COMPARE_RESULTS) then
    COMPARE_RESULTS -i $date >>& $results &; wait
    @ return = $status
    if ($return != 0) then
        echo 'Status: '$return' COMPARE_RESULTS exited with ERROR'
    endif
    echo ' ' >>& $results
else
    echo "Can't find COMPARE_RESULTS - check your verification directory"
    exit 1
endif

echo $version: Verification completed "`date`" >>& $results
```



## F. RUN\_VERIFICATION Script

### RUN\_VERIFICATION script

```
#!/bin/csh
#   RUN_VERIFICATION
#   Script for execution of verification problems

echo 'Verification Runs for the FEHMN Application'
date
echo ' '

@ paths = 0

# Determine if executable and verification directory are defined
if (!($?XFEHMN) || !($?VER_DIR)) then

# Look for paths on the command line
    if ($#argv > 0 && $1 == "-p") then
        @ paths = 1
        shift argv
        setenv XFEHMN $1
        shift argv
        setenv VER_DIR $1
        shift argv

# Look in PATHS file
    else if (-f PATHS) then
        @ paths = 2
        source PATHS

    else
        echo 'PATHs to the FEHMN executable and verification directory must be set.'
        echo 'They can be entered as command line arguments or put in a file'
        echo 'called PATHS in the current directory.'
        echo ' '
        echo 'usage:RUN_VERIFICATION [-p xfehm_n_path verification_dir] [problem1 problem2 ...]'
        exit 1
    endif
endif

@ flag = 0

# Verify that XFEHMN is executable
if (!(-x $XFEHMN)) then
    echo '$XFEHMN' does not exist or is not an executable file.'
    @ flag = 1
endif

# Verify that VER_DIR is a directory
if (!(-d $VER_DIR)) then
    echo '$VER_DIR' is not a directory.'
    @ flag = 1
endif
```

# RUN\_VERIFICATION script (continued)

```

if ($flag != 0) then
    if ($paths == 2) then
        echo 'Check your PATHS file'
    else if ($paths == 1) then
        echo 'Check your command line input'
    else
        echo 'Check FEHM_VVSECT input'
    endif
    exit 1
endif

# If no problems are specified execute all tests
if ($#argv == 0) then
    foreach problem (avdonin dissolution doe5a dryout dual \
        fracture_transport heat2d heat3d henrys_law \
        infiltration multi_solute ramey sorption theis toronyi \
        transport3D vapor_extraction)
        echo '***** BEGIN '$problem' *****'
        if (-e $VER_DIR/$problem && -d $problem) then
            setenv problem $problem
            echo 'cd '$problem'; '$VER_DIR'/'$problem'/run'$problem'; wait'
            cd $problem; $VER_DIR/$problem/run$problem; wait
            cd ..
        else
            echo $VER_DIR/'$problem' or '$problem' does not exist'
        endif
        echo '***** END '$problem' *****'
        echo ' '
    end

# Else execute just the specified problems
else
    while ($#argv > 0)
        echo '***** BEGIN '$1' *****'
        if (-e $VER_DIR/$1 && -d $1) then
            setenv problem $1
            echo 'cd '$1'; '$VER_DIR'/'$1'/run'$1'; wait'
            cd $1; $VER_DIR/$1/run$1; wait; cd ..
        else
            echo $VER_DIR/'$1' or '$1' does not exist'
        endif
        echo '***** END '$1' *****'
        echo ' '
        shift argv
    end
endif

echo 'Verification Runs Completed'
date

```

## G. COMPARE\_RESULTS Script

### COMPARE\_RESULTS script

```
#!/bin/csh
# COMPARE_RESULTS
# Script for generating comparison of results and summary

echo 'Compare Results for the FEHMN Application Verification Runs'
date
echo ' '

@ path_flag = 0

# Determine if verification directory and executables are defined

if (!($?VER_DIR)) then

# Look for PATHS on the command line
    if ($#argv >= 2 && $1 == "-p") then
        @ path_flag = 1
        shift argv
        setenv VER_DIR $1
        shift argv

# or Look in PATHS file
    else if (-f PATHS) then
        @ path_flag = 2
        source PATHS

    else

        echo 'PATHS to the verification directory must be set.'
        echo 'It can be entered as a command line argument or put in a file'
        echo 'called PATHS in the current directory.'
        echo ' '
        echo 'usage:COMPARE_RESULTS [-p verification_dir] [-i date_id] [problem1 problem2 ...]'
        exit 1

    endif

endif

# Verify that VER_DIR is a directory
if (!(-d $VER_DIR)) then
    echo $VER_DIR' is not a directory.'

    if ($path_flag == 2) then
        echo 'Check your PATHS file'
    else ($path_flag == 1) then
        echo 'Check your command line input'
```

**COMPARE\_RESULTS script (continued)**

```
    else
        echo 'Check FEHM_VVSECT input'
    endif

    exit 1

endif

# Define verification executables
setenv COMPARE      $VER_DIR/COMPARE
setenv COMPARET     $VER_DIR/COMPARET
setenv SUMMARIZE    $VER_DIR/SUMMARIZE

@ flag = 0

# Verify that verification executables exist / can be executed
if (! -x $COMPARE) then
    echo $COMPARE' does not exist or is not an executable file.'
    @ flag = 1
endif
if (! -x $COMPARET) then
    echo $COMPARET' does not exist or is not an executable file.'
    @ flag = 1
endif
if (! -x $SUMMARIZE) then
    echo $SUMMARIZE' does not exist or is not an executable file.'
    @ flag = 1
endif

if ($flag != 0) then
    echo 'Check your verification directory: '$VER_DIR
    exit 1
endif

if ($#argv == 0 || $1 != "-r") then
    set date=`date +%y%m%d`
else if ($1 == "-r") then
    shift argv
    set date = $1
    shift argv
endif

set summary = SUMMARY_RPT.$date
if (-e $summary) then
    mv $summary $summary.old
endif

echo 'SUMMARY of FEHM COMPARISON TESTS '$date > $summary
echo ' ' >> $summary
```

# COMPARE\_RESULTS script (continued)

```
# If no problems are specified execute all tests
if ($#argv == 0) then

    foreach problem (avdonin dissolution doe5a dryout dual \
        fracture_transport heat2d heat3d henrys_law \
        infiltration multi_solute ramey sorption this \
        toronyi transport3D vapor_extraction)
        echo '***** BEGIN '$problem' *****'
        echo '***** BEGIN '$problem' *****' >> $summary
        if (-e $VER_DIR/$problem && -d $problem) then
            setenv problem $problem
            echo 'cd '$problem'; comp'$problem' '$date
            cd $problem; $VER_DIR/$problem/comp$problem $date; cd ..
            cat $problem/summary.$date >> $summary
        else
            echo $VER_DIR/'$problem' or '$problem' does not exist'
        endif
        echo '***** END '$problem' *****'
        echo '***** END '$problem' *****' >> $summary
        echo ' '
        echo ' ' >> $summary
    end

# Else compare just the specified problems
else

    while ($#argv > 0)
        echo '***** BEGIN '$1' *****'
        echo '***** BEGIN '$1' *****' >> $summary
        if (-e $VER_DIR/$1 && -d $1) then
            setenv problem $1
            echo 'cd '$1'; comp'$1' '$date
            cd $1; $VER_DIR/$1/comp$1 $date; cd ..
            cat $1/summary.$date >> $summary
        else
            echo $VER_DIR/'$1' or '$1' does not exist'
        endif
        echo '***** END '$1' *****'
        echo '***** END '$1' *****' >> $summary
        echo ' '
        echo ' ' >> $summary
        shift argv
    end

endif

echo 'End Compare Results for the FEHMN Application Verification Runs'
date
```

## H. Example of Problem Execution Script

### runavdonin script

```
#!/bin/csh
#      runavdonin

if (! -d input) then
    set INPUT = $VER_DIR/$problem/input
    rm -f input
    ln -s $INPUT input
endif

foreach geom (84 400 800)
    echo 'sed s/base/'$geom'/ input/avdonin.files > fehmn.files'
    sed s/base/$geom/ input/avdonin.files > fehmn.files
    echo 'nice '$XFEHMN' &; wait'
    nice $XFEHMN &; wait
end
rm fehmn*
```

## I. Example of Run Comparison Script

### compavdonin script

```
#!/bin/csh
#   Comparisons for avdonin problem

if ($#argv == 0) then
    set ID = `date +%y%m%d`
else
    set ID = $1
endif

if (-e summary.$ID) then
    mv summary.$ID summary.$ID.old
endif
echo 'Summary file named: summary.$ID'

if (! -d input) then
    set INPUT = $VER_DIR/$problem/input
    rm -f input
    ln -s $INPUT input
endif

foreach type (history contour)

    foreach geom (84 400 800)
        echo 'compare '$geom' '$type
        sed s/base/$geom/ input/avdonin.comparein.$type > comparein
        nice $COMPARE &; wait
    end

    if ($type == history) then
        sed s/param/time/ input/avdonin.summary > summarize
    else if ($type == contour) then
        sed s/param/pos/ input/avdonin.summary > summarize
    endif
    nice $SUMMARIZE >> summary.$ID; wait

end
rm comparein* summarize
```

## J. Execution Log

### Example of execution log

FEHM 01.00: Verification started Tue Jul 23 13:17:44 MDT 1996

Verification Runs for the FEHMN Application  
Tue Jul 23 13:17:45 MDT 1996

\*\*\*\*\* BEGIN avdonin \*\*\*\*\*

cd avdonin; /home/fehm/verification/avdonin/runavdonin; wait

sed s/base/84/ input/avdonin.files > fehm.files

nice /home/fehm/bin/xfehm.95-05-01p-sun4 &; wait

[1] 25919

[1] Done /home/fehm/bin/xfehm.95-05-01p-sun4

sed s/base/400/ input/avdonin.files > fehm.files

nice /home/fehm/bin/xfehm.95-05-01p-sun4 &; wait

[1] 25923

[1] Done /home/fehm/bin/xfehm.95-05-01p-sun4

sed s/base/800/ input/avdonin.files > fehm.files

nice /home/fehm/bin/xfehm.95-05-01p-sun4 &; wait

[1] 25929

[1] Done /home/fehm/bin/xfehm.95-05-01p-sun4

\*\*\*\*\* END avdonin \*\*\*\*\*

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\*\*\*\*\* BEGIN vapor\_extraction \*\*\*\*\*

cd vapor\_extraction; /home/fehm/verification/vapor\_extraction/runvapor\_extraction; wait

sed s/case/iso/ input/vapextract.files > fehm.files

nice /home/fehm/bin/xfehm.95-05-01p-sun4 &; wait

[1] 26145

[1] Done /home/fehm/bin/xfehm.95-05-01p-sun4

sed s/case/aniso/ input/vapextract.files > fehm.files

nice /home/fehm/bin/xfehm.95-05-01p-sun4 &; wait

[1] 26147

[1] Done /home/fehm/bin/xfehm.95-05-01p-sun4

\*\*\*\*\* END vapor\_extraction \*\*\*\*\*

Verification Runs Completed  
Tue Jul 23 20:53:53 MDT 1996



**Example of execution log (continued)**

Compare Results for the FEHMN Application Verification Runs  
Tue Jul 23 20:53:54 MDT 1996

```
***** BEGIN avdonin *****
cd avdonin; compavdonin 950803
Summary file named: summary.950803
compare 84 history
[1] 26156
[1] Done          /home/fehm/verification/COMPARE
compare 400 history
[1] 26158
[1] Done          /home/fehm/verification/COMPARE
compare 800 history
[1] 26160
[1] Done          /home/fehm/verification/COMPARE
compare 84 contour
[1] 26164
[1] Done          /home/fehm/verification/COMPARE
compare 400 contour
[1] 26166
[1] Done          /home/fehm/verification/COMPARE
compare 800 contour
[1] 26168
[1] Done          /home/fehm/verification/COMPARE
***** END avdonin *****

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***** BEGIN vapor_extraction *****
cd vapor_extraction; compvapor_extraction 950803
Summary file named: summary.950803
compare iso contour
[1] 26587
[1] Done          /home/fehm/verification/COMPARE
compare aniso contour
[1] 26590
[1] Done          /home/fehm/verification/COMPARE
***** END vapor_extraction *****
```

End Compare Results for the FEHMN Application Verification Runs  
Tue Jul 23 20:56:34 MDT 1996

FEHM 01.00: Verification completed Tue Jul 23 20:56:34 MDT 1996

## K. Summary Report

### Example of summary report

#### SUMMARY of FEHM COMPARISON TESTS 960723

\*\*\*\*\* BEGIN avdonin \*\*\*\*\*

Avdonin Radial Heat and Mass Transfer Problem Comparison of Model and Analytical Solution --  
Temperature vs Time  
At R coordinate 37.5000

Test Case	Maximum Error	Maximum % Error	RMS Error
84 nodes	1.262	0.7776	0.2169E-03
400 nodes	0.4060	0.2487	0.6973E-04
800 nodes	0.3899	0.2384	0.6742E-04

Avdonin Radial Heat and Mass Transfer Problem Comparison of Model and Analytical Solution --  
Temperature vs Position  
At Time 0.100000E+10

Test Case	Maximum Error	Maximum % Error	RMS Error
84 nodes	0.5230	0.3237	0.1745E-03
400 nodes	0.2815	0.1744	0.3416E-04
800 nodes	0.2815	0.1744	0.2213E-04

\*\*\*\*\* END avdonin \*\*\*\*\*

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\*\*\*\*\* BEGIN vapor\_extraction \*\*\*\*\*

Vapor Extraction - Vapor Pressure vs Position

Test Case	Maximum Error	Maximum % Error	RMS Error
anisotropic	0.3066E-02	3.311	0.1436E-0
isotropic	0.1983E-02	2.195	0.8838E-04

\*\*\*\*\* END vapor\_extraction \*\*\*\*\*

