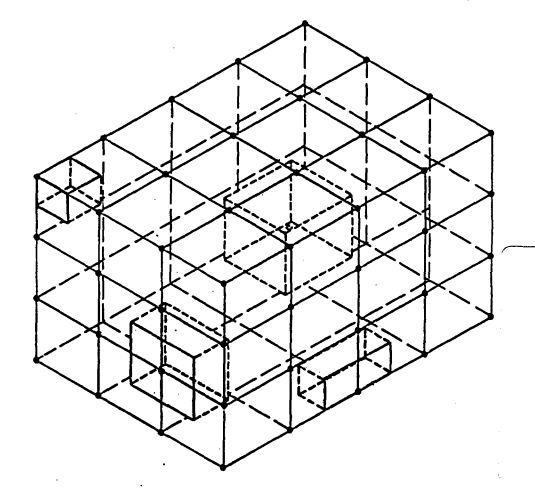
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HST3D: A COMPUTER CODE FOR SIMULATION OF HEAT AND SOLUTE TRANSPORT IN THREE-DIMENSIONAL GROUND-WATER FLOW SYSTEMS

# U.S. GEOLOGICAL SURVEY



Water-Resources Investigations Report 86-4095

UNITED STATES DEPARTMENT OF THE INTERIOR

DONALD PAUL HODEL, Secretary

GEOLOGICAL SURVEY

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By Kenneth L. Kipp, Jr.

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U.S. GEOLOGICAL SURVEY

Water-Resources Investigations Report 86-4095



Denver, Colorado 1987

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# CONVERSION FACTORS

The HST3D simulator program performs calculations in metric units. `However, it will accept input and produce output in inch-pound units. The . conversion factors are listed below:

Multiply	By	To obtain
ک kilogram (kg)	2.204622	
meter (m)		pound (1b)
millimeter(mm)	3.280840	foot (ft)
second (s)	$3.937008 \times 10^{-2}$	inch (in.)
	$1.157407 \times 10^{-5}$	day (d)
degree Celsius (°C)	$T(^{\circ}F) = 1.8T(^{\circ}C) + 32$	degree Fahrenheit (°F)
Kelvin (K)	$T(^{\circ}F) = 1.8T(K) - 459.67$	degree Fahrenheit (°F)
Joule (J) or Watt- second (W-s)	9.478170 × 10 <sup>-4</sup>	British Thermal Unit (BTU)
square meter (m <sup>2</sup> )	10.76391	square foot (ft <sup>2</sup> )
cubic meter (m <sup>3</sup> )	35.31466	cubic foot (ft <sup>3</sup> )
meter-second (m-s)	$3.797267 \times 10^{-5}$	foot-day (ft-d)
Pascal (Pa)	$1.450377 \times 10^{-4}$	pound per square inch (psi)
meter per second (m/s)	$2.834646 \times 10^{5}$	foot per day (ft/d)
square meter per second (m <sup>2</sup> /s)	$9.300018 \times 10^{5}$	square foot per day (ft <sup>2</sup> /d)
<pre>cubic meter per second   (m<sup>3</sup>/s)</pre>	$3.051187 \times 10^6$	cubic foot per day (ft <sup>3</sup> /d)
liter per second (2/s)	$3.051187 \times 10^3$	cubic foot per day (ft <sup>3</sup> /d)
kilogram per second (kg/s)	$1.904794 \times 10^{5}$	pound per day (1b/d)
Pascal per second (Pa/s)	12.53126	<pre>pound per square inch     per day (lb/in<sup>2</sup>/d)</pre>
cubic meter per cubic meter-second (m <sup>3</sup> /m <sup>3</sup> -s)	8.6400 × 10 <sup>4</sup>	cubic foot per cubic foot-day (ft <sup>3</sup> /ft <sup>3</sup> -d)
kilogram per cubic meter (kg/m <sup>3</sup> )	$6.242797 \times 10^{-2}$	pound per cubic foot (lb/ft <sup>3</sup> ) <sup>1</sup>
Watt per cubic meter (W/m <sup>3</sup> )	9.662109 × $10^{-2}$	British Thermal Unit per hour-cubic foot (BTU/h-ft <sup>3</sup> )
Joule per kilogram (J/kg)	$4.299226 \times 10^{-4}$	British Thermal Unit per pound (BTU/1b)
Joule per kilogram (J/kg)	0.3345526	foot-pound force per pound mass (ft-lbf/lbm)
cubic meter per kilogram (m <sup>3</sup> /kg)	16.01846	cubic foot per pound (ft <sup>3</sup> /1b)
cubic meter per square meter-second (m <sup>3</sup> /m <sup>2</sup> -s)	$2.834646 \times 10^{5}$	cubic foot per square foot-day (ft <sup>3</sup> /ft <sup>2</sup> -d)
Watt per square meter (W/m <sup>2</sup> )	0.3169983	British Thermal Unit per hour-square foot (BTU/h-ft <sup>2</sup> )
kilogram per square meter-second (kg/m <sup>2</sup> -s)	1.769611 × 10 <sup>€</sup>	pound per square foot-day (lb/ft <sup>2</sup> -d)

9.300013 × 10<sup>5</sup> cubic meter per meter-second  $(m^3/m-s)$ kilogram per meter-second 1,000 (kg/m-s)Joule per kilogram-meter  $1.310404 \times 10^{-4}$ (J/kg-m)Watt per meter-degree 13.86941 Celsius (W/m-°C) Watt per square meter-0.1761102 degree Celsius (W/m<sup>2</sup>-°C) Joule per kilogram- $2.388459 \times 10^{-4}$ degree Celsius (J/kg-°C) Joule per cubic meter-degree  $1.491066 \times 10^{-5}$ Celsius (J/m<sup>3</sup>-°C) cubic meter per second-meter-  $6.412138 \times 10^9$ Pascal  $(m^3/s-m-Pa)$ 

A weight density rather than a mass density.
 Not inch-pound but common usage.

cubic foot per foot-day {
 (ft<sup>3</sup>/ft-d)
 centipoise (cP)<sup>2</sup>

British Thermal Unit per pound-foot (BTU/1b-ft) British Thermal Unit • per foot-hour-degree Fahrenheit (BTU/ft-h-°F) British Thermal Unit per hour-square foot-degree Fahrenheit  $(BTU/h-f\iota^2-{}^{\circ}F)$ British Thermal Unit jer pound-degree Fahrenheit (BTU/15-°F) British Thermal Unit per cubic foot-degree Fahrenheit  $(BTU/ft^3-{}^{\circ}F)$ cubic foot per dayfoot-pound-square inch (ft<sup>3</sup>/d-ft-psi)

#### HST3D: A COMPUTER CODE FOR SIMULATION OF HEAT AND SOLUTE TRANSPORT

## IN THREE-DIMENSIONAL GROUND-WATER FLOW SYSTEMS

By Kenneth L. Kipp Jr.

#### ABSTRACT

The Heat- and Solute-Transport Program (HST3D) simulates ground-water flow and associated heat and solute transport in three dimensions. The HST3D program may be used for analysis of problems such as those related to subsurface-waste injection, landfill leaching, saltwater intrusion, freshwater recharge and recovery, radioactive-waste disposal, hot-water geothermal systems, and subsurface-energy storage. The three governing equations are coupled through the interstitial pore velocity, the dependence of the fluid density on pressure, temperature, and solute-mass fraction, and the dependence of the fluid viscosity on temperature and solute-mass fraction. The solutetransport equation is for only a single, solute species with possible linearequilibrium sorption and linear decay. Finite-difference techniques are used to discretize the governing equations using a point-distributed grid. The flow-, heat- and solute-transport equations are solved, in turn, after a partial Gauss-reduction scheme is used to modify them. The modified equations are more tightly coupled and have better stability for the numerical solutions.

The basic source-sink term represents wells. A complex well-flow model may be used to simulate specified flow rate and pressure conditions at the land surface or within the aquifer, with or without pressure and flow-rate constraints. Boundary-condition types offered include specified value, specified flux, leakage, heat conduction, an approximate free surface, and two types of aquifer-influence functions. All boundary conditions can be functions of time.

Two techniques are available for solution of the finite-difference matrix equations. One technique is a direct-elimination solver, using equations reordered by alternating diagonal planes. The other technique is an iterative solver, using two-line successive overrelaxation. A restart option is available for storing intermediate results and restarting the simulation at an intermediate time with modified boundary conditions. This feature also can be used as protection against computer-system failure.

Data input and output may be in metric (SI) units or inch-pound units. Output may include tables of dependent variables and parameters, zoned-contour maps, and plots of the dependent variables versus time. The HST3D program is a descendant of the Survey Waste Injection Program (SWIP) written for the U.S. Geological Survey under contract.

#### 1. INTRODUCTION

### 1.1. OVERVIEW OF THE SIMULATOR

The computer program (HST3D) described in this report simulates heat and solute transport in three-dimensional saturated ground-water flow systems. The equations that are solved numerically are: (1) The saturated ground-water flow equation, formed from the combination of the conservation of total-fluid mass and Darcy's Law for flow in porous media; (2) the heat-transport equation from the conservation of enthalpy for the fluid and porous medium; and (3) the solute-transport equation from the conservation of mass for a single-solute species, that may decay and may adsorb onto the porous medium. These three equations are coupled through the dependence of advective transport on the interstitial fluid-velocity field, the dependence of fluid viscosity on temperature and solute concentration, and the dependence of fluid density on pressure, temperature, and solute concentration.

Numerical solutions are obtained for each of the dependent variables: pressure, temperature, and mass fraction (solute concentration) in turn, using a set of modified equations that more directly link the original equations through the velocity-, density-, and viscosity-coupling terms. Finitedifference techniques are used for the spatial and temporal discretization of the equations. When supplied with appropriate boundary and initial conditions and system-parameter distributions, simulation calculations can be performed to evaluate a wide variety of heat- and solute-transport situations.

The computer code (HST3D) described in this documentation is a descendant of a computer code for calculating the effects of liquid-waste disposal into deep, saline aquifers, developed by INTERCOMP Resource Development and Engineering Inc. 1976) for the U.S. Geological Survey and revised by INTERA Environmental Consultants Inc. (1979). The parent code, known as the Survey Waste Injection Program (SWIP), has been completely rewritten with many major and minor modifications, improvements, and correction of several errors. Features included in HST3D are briefly described as follows:

- Specified-value and specified-flux boundary conditions are independent of each other and independent of the well or aquifer-influence-function boundary conditions. The boundary conditions also may vary with time.
- 2. Specified heat- and solute-flux boundary conditions are available.
- 3. The leakage boundary conditions are generalized and a riverleakage boundary condition is available.
- 4. Porous-medium thermal properties, dispersivity, and compressibility, may have spatial variation defined by zones.
- 5. A point-distributed, finite-difference grid is employed, rather than a cell- or block-centered grid, for less truncation error and easier incorporation of boundary conditions.
- 6. The heat-conduction boundary condition is generalized to apply to any cell face.
- 7. Global-flow, and heat- and solute-balance calculations are performed including flux calculations through specified pressure, temperature, and mass-fraction boundaries.
- 8. A robust algorithm for the computation of the optimum overrelaxation factor for the two-line, successive-overrelaxation, matrix-solution method is used, with a convergence criterion that includes the matrix spectral-radius estimate.
- 9. The code is organized for a logical flow of calculation and a modular structure.
- 10. The code length is about 12,000 lines, using FORTRAN 77 language constructs for cleaner, more efficient coding than possible with FORTRAN 66. However, clarity has not been sacrificed for ultimate efficiency.
- Comments have been included liberally for ease of understanding the program.
- 12. All arrays with lengths depending on the size of the problem are in two variably-partitioned arrays, integer and real, to facilitate double-precision arithmetic.

- 13. Arrays required for thermal or solute calculations exclusively are eliminated if only one of these transported quantities is being simulated, which results in a considerable decrease in computer storage.
- 14. Arrays used for a specific type of boundary condition or source-sink condition are dimensioned only to the length required.
- 15. The allocation of space for the direct-equation solver is explicitly determined during array-space allocation, rather than estimated.
- 16. Logical variables are used to control the flow of program execution for ease of option selection.
- 17. The input file is in free-format to facilitate input from terminals.
- 18. The input file is organized into logical groups for parameter specifications.
- 19. User comments can be freely incorporated into the input file for rapid identification of the data. An input-file form is available which the user can fill out at the terminal for a given simulation.
- 20. A read-echo file may be written to aid in locating errors in the data-input file.
- 21. Character plots of the porous-media zones may be created on the output file to facilitate checking the zonation.
- 22. Although the internal calculations of the program are performed in metric units, the input and output can be chosen to be in inch-pound units.
- 23. The output material is made easily understandable by avoiding variable names, by logical grouping on the page, and by including supplementary information.
- 24. Error tests are included to catch likely mistakes in data input.
- 25. Error messages are printed explicity rather than as code numbers.

- 26. There is no limit on the number of plots that can be created. The number of calculated points in time per plot is limited to three times the total number of grid points, while the number of observed points in time is limited to two times the number of grid points. The user can select every nth point to be plotted, if this number is limiting.
- 27. The solute concentration can be chosen to be the mass fraction or a scaled mass fraction that ranges from 0 to 1. This choice was available in the SWIP code, but the user was not clearly made aware of which option was selected.
- 28. Two types of restart option are available: a periodic check-point dump for protection against computer-system failure, and a specific dump for user review and possible modification of parameters.
- 29. Map-contour intervals can be automatically determined to be a multiple of 2, 5, or 10, and the contour zones are "zebra striped" for easier reading.
- 30. Initial-pressure conditions can be specified to be other than hydrostatic. For example, an initial water-table configuration can be used.
- 31. The precipitation-infiltration option is contained in the distributed flux-boundary conditions.
- 32. The conductive-heat-loss to overburden and underburden is a general, heat-transfer calculation, applicable to any cell face in the region.
- 33. The well-riser, heat-transfer calculation is based on heat transfer from a known-temperature, cylindrical boundary, and higher order assymptotic expansions have been used.
- 34. The well-riser calculation has been formulated to solve the total-energy and momentum balance equations simultaneously, using the Bulirsch-Stoer algorithm for integration of the ordinary differential equations.
- 35. The well-bore equations are implicitly coupled to the system equations for cases of cylindrical geometry.

- 3b. The well-datum pressure and the well-flow rate allocation calculations may be performed iteratively in conjunction with the solution of the flow equation, or explicitly.
- 37. The full nine-component, or an approximate three-component, dispersion-coefficient tensor may be used for cross-dispersive flux calculations.

The purpose of simulation modeling the transport of heat and solute in pround-water flow systems is to gain a quantitative understanding of how the sources and sinks, the boundary conditions, and the aquifer parameters interact to cause ground-water flow patterns and consequent thermal- and solute-concentration movement in a system under investigation. Of particular interest are the magnitudes of concentrations and discharges at interfaces with the environment, for example, in cases of aquifer contamination. Naturally, the quality or degree of realism of a given simulation is strongly dependent on the quantity and quality of the parameter distribution, boundarycondition, and source-sink data. Acquiring this data can be a major task of the modeling project.

## 1.2. APPLICABILITY AND LIMITATIONS

The HST3D code is suitable for simulating ground-water flow and the associated heat and solute transport, in saturated, three-dimensional flow systems with variable density and viscosity. As such, the code is applicable to the study of waste injection into saline aquifers, landfill-contaminant movement, seawater intrusion in coastal regions, brine disposal, fresh-water storage in saline aquifers, heat storage in aquifers, liquid-phase geothermal systems, and similar transport situations. If desired, only the ground-water flow or only the heat- or the solute-transport equation may be solved in conjunction with ground-water flow. Three-dimensional cartesian or axisymmetric, cylindrical-coordinate systems are available.

The primary limitation of this code results from the use of finitediffc-ence techniques for the spatial- and temporal-derivative approximations.

Where longitudinal and transverse dispersivities may be small, cell sizes will need to be small to minimize numerical dispersion or oscillation. Furthermore, if the region of solute movement is somewhat convoluted and threedimensional, the projection of nodal lines from regions of high-nodal density will cause more nodes than are needed to appear in other regions. These two factors can combine to cause an excessive number of nodes to be involved for a given simulation, thus making the simulation prohibitively expensive because of computer-storage and computation-time requirements. In such cases, a simple model of the system, useful for investigating mechanisms and testing hypotheses, may be all that is practical.

Another limitation results from a phenomenon called grid-orientation effect (Aziz and Settari, 1979, p. 332), whereby numerical simulations of miscible displacement converge to two separate solutions, as the mesh size is refined, depending on whether the major velocity vectors are parallel to one of the coordinate directions or are diagonally oriented. The effect is more pronounced for conditions of little dispersion or piston-like displacement of the solute, and for conditions of the viscosity of the displacing fluid much less than the viscosity of the displaced fluid. The effect virtually is absent if the two viscosities are nearly equal, or if the dispersion coefficient is large. The primary cause of the grid-orientation effect appears to be the use of a seven-point difference formula for the threedimensional-flow and solute-transport equations, because this formula restricts transport in the diagonal directions. Use of a grid where the major velocity vectors are oriented parallel to one of the coordinate directions, has been found to give more realistic simulation results (Aziz and Settari, 1979, p. 336). To completely eliminate this problem, a higher-order differencing scheme, or curvilinear coordinates need to be used, but these modifications are beyond the scope of the present version of HST3D.

There is a limitation on which boundary conditions can be used with a tilted coordinate system. The free surface and leakage boundary conditions require that the z-axis be oriented in the vertical direction.

A limitation that is secondary for most ground-water flow and transport modeling is that two types of transport phenomena exist that this type of numerical simulation has difficulty in representing quantitatively. The first phenomenon, viscous-fingering instabilities, may occur during the displacement of a resident fluid by an injected fluid with significantly less viscosity. The injected fluid forms channels or fingers through the resident fluid, as described by Aronofsky (1952), Saffman and Taylor (1958), and Sheidegger (1960). The second phenomenon may occur in the situation where a fluid of greater density overlies one of lesser density. Rayleigh-Taylor convective cells are formed that mix the two fluids (Wooding, 1959). Numerical simulation tends to predict these transport instabilities later than they occur in laboratory-scale experiments. When perturbations are present to initiate the instabilities, the general magnitudes often are calculated to be less than those that actually occur (Scheidegger and Johnson, 1963; and Dougherty, 1963). However, laboratory-scale viscous fingering and convectivecell formation may be much more unstable than the corresponding field-scale phenomenon, because of the smaller dispersivity at the laboratory scale. Therefore, at the field scale, numerical simulation may not be so much in error in representing these instabilities. Nevertheless, these limitations need to be kept in mind when simulating fluid flow with large viscosity or density contrasts.

Another secondary limitation is that this is a rather general computer code. The variety of discretization, boundary-condition, and source-sink options make this code not as computationally efficient as a simulation code designed specifically for a given system being investigated. This limitation is compensated by the ability of the HST3D simulator to represent a wide variety of physical situations.

#### 1.3. PURPOSE AND SCOPE

The purpose of this documentation is to provide the user with information on the theory, assumptions, and equations being numerically solved, the numerical-solution methods employed, and the various program options available. The sets of verification test problems are presented and two example problems are described in detail with input and output files. Sections on the code organization, input information, and output information, as well as a list of variable-definitions and a cross-reference map are provided. The documentation is intended to be sufficiently complete and understandable so the user easily can obtain successful simulations, diagnose most computational problems, develop remedies, and incorporate minor program additions or modifications to suit specific modeling needs.

Each release of the HST3D program code is identified by a release number. This documentation is for release 1.0, and this number will change as modifications, corrections, and additions are made to the program. Updates to the documentation will be keyed to the release number.

#### 1.4. ACKNOWLEDGMENTS

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#### 2. THEORY

### 2.1. FLOW AND TRANSPORT EQUATIONS

Derivation of the saturated ground-water flow and heat- and solutetransport equations solved by this program can be found in references such as Bear (1972) or Huyakorn and Pinder (1983). Only the assumptions leading to these equations will be presented here. Explanations of the notation will appear after the first usage. A complete table of notation appears in chapter 9. In the report, all variables will be given with metric (SI) units of measure.

## 2.1.1. Ground-Water Flow Equation

The partial-differential equation of ground-water flow is based on the following assumptions:

- Ground water fully saturates the porous medium within the region of ground-water flow.
- Ground-water flow is described by Darcy's Law.
- The porous medium is compressible.
- The fluid is compressible.
- The porosity and permeability are functions of space.
- The coordinate system is chosen to be aligned with the principal directions of the permeability tensor so that this tensor is diagonal for anisotropic media.
- The coordinate system is orthogonal as are the principal directions of the permeability tensor.
- The coordinate system is right-handed with the z-axis pointing vertically upward.
- The fluid viscosity is a function of space and time through dependence on temperature and solute concentration.
- Density-gradient diffusive fluxes of the bulk fluid are neglected relative to advective-mass fluxes.

- Dispersive-mass fluxes of the bulk fluid from spatial-velocity fluctuations are not included.
- Contributions to the total fluid-mass balance from pure-solutemass sources within the region are not included.

Pressure is chosen as the dependent variable for fluid flow, because no potentiometric-head function exists for density fields that depend on temperature and solute concentration. All pressures denoted by p are expressed relative to atmospheric pressure. Absolute pressures are denoted by  $\hat{p}$ . The flow equation is based on the conservation of total fluid mass in a volume element, coupled with Darcy's Law for flow through a porous medium. Thus:

$$\frac{\partial(\epsilon\rho)}{\partial t} = \nabla \cdot \rho \frac{\underline{k}}{\mu} (\nabla p + \rho g) + q \rho^* ; \qquad (2.1.1.1a)$$

where

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p is the fluid pressure (Pa);
t is the time (s);
ɛ is the effective porosity (-);
p is the fluid density (kg/m<sup>3</sup>);
p* is the density of a fluid source (kg/m<sup>3</sup>);
k is the porous-medium permeability tensor (m<sup>2</sup>);
µ is the fluid viscosity (kg/m-s);
g is the gravitational constant (m/s<sup>2</sup>); and
q is the fluid-source flow-rate intensity (m<sup>3</sup>/m<sup>3</sup>-s); (positive
is into the region).
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Equation 2.1.1.1a relates the rate of change of total mass in the fluid phase to net fluid-inflow rate, and source fluid-and-solute flow rate. Note that the density of the fluid source is  $\rho^*$  for q>0, and  $\rho$  for q<0.

The interstitial or pore velocity, v is obtained from Darcy's Law as:

(2.1.1.1b)

$$\underline{v} = - \frac{\underline{k}}{\epsilon \mu} (\nabla p + \rho g);$$

where

v is the interstitial-velocity vector (m/s).

#### 2.1.2. Heat-Transport Equation

The thermal-energy-balance equation, used for heat transport, is based on the following assumptions:

- Fluid kinetic energy is negligible.
- Thermal-dispersive transport takes place with a mechanism analogous to solute-dispersive transport.
- Thermal conduction occurs through the fluid and porous medium in parallel.
- Radiant-energy transfer is neglected.
- Thermal effects of chemical reactions are neglected.
- Changes in gravitational energy from diffusive and dispersive fluxes of solute species are neglected.
- Heating from viscous dissipation is neglected.
- Heat capacities are not a function of temperature or solute concentration.
- Thermal conductivities are not functions of temperature or solute concentration.
- Thermal equilibrium exists between the fluid and solid phases.
- Energy transport by a diffusive flux of solute is neglected.
- Only a single fluid phase exists.
- Pressure equilibrium exists between the fluid and porous-medium phases.
- Changes in fluid enthalpy with pressure, that is, pressure volume work, reversible work, or flow work, as a parcel of fluid moves are neglected.

- The velocity of the porous medium during compression or expansion 15 neglected.
- Enthalpy dependence on solute concentration is accounted for by a heat-capacity adjustment.
- The thermal expansion of the porous medium is neglected.

The energy equation is based upon the conservation of enthalpy in both the fluid and solid or porous-medium phases of a volume of the region. Enthalpy is a derived property containing both internal energy and flow energy. Temperature is the dependent variable. Thus:

 $\frac{\partial}{\partial t} (\epsilon \rho c_{f} + (1 - \epsilon) \rho_{s} c_{s})T = \nabla \cdot (\epsilon K_{f} + (1 - \epsilon) K_{s}) \underline{I} \nabla T$ 

+  $\nabla \cdot \varepsilon \underline{D}_{\mu} \nabla T - \nabla \cdot \varepsilon \rho c_{\mu} \underline{v} T$ 

+  $q_{H}$  +  $qp*c_{f}T*$ ; (2.1.2.1)

where

T is the fluid and porous-medium temperature (°C);
T\* is the temperature of the fluid source (°C);
ρ<sub>s</sub> is the density of the solid phase (kg/m<sup>3</sup>);
c<sub>f</sub> is the heat capacity of the fluid phase at constant pressure (J/kg-°C);
c<sub>s</sub> is the heat capacity of the solid phase at constant pressure (J/kg-°C);
K<sub>f</sub> is the thermal conductivity of the fluid phase (W/m-°C);
K<sub>s</sub> is the thermal conductivity of the solid phase (W/m-°C);
E<sub>H</sub> is the thermo-mechanical dispersion tensor (W/m-°C);
q<sub>H</sub> is the heat-source rate intensity (W/m<sup>3</sup>); and
L is the identity matrix of rank 3 (-).

Equation 2.1.2.1 relates the rate of change of fluid and porous-medium enthalpy to the net conductive-enthalpy flux, to the net dispersive enthalpy flux, to the net advective-enthalpy flux, to the heat source, and to the fluid source at a given temperature. It is written for a unit volume of fluid and solid phase together; that is, a unit volume of saturated, porous medium. Heat is injected at temperature, T\*, and density,  $\rho$ \*, by a fluid source; but heat is withdrawn at temperature, T, and density,  $\rho$  by a fluid sink. A detailed derivation of equation 2.1.2.1 is given in Faust and Mercer (1977).

#### 2.1.3. Solute-Transport Equation

The equation for conservation of a single solute species is based on the following assumptions:

- Thermal diffusion is neglected.
- Pressure diffusion is neglected.
- Solute transport by local, interstitial, velocity-field fluctuations and mixing at pore junctions is described by a hydrodynamicdispersion coefficient.
- Forced diffusion by gravitational, electrical, and other fields is neglected.
- The only reaction mechanism is linear decay or disappearance of solute.
- The only solute, porous-medium, interaction mechanism is linearequilibrium sorption.
- No pure solute sources occur in the fluid or solid phases.

The solute mass fraction is taken to be the dependent variable because the density field is variable. It is an amount per unit mass of fluid, that is, a mass-based concentration. The more widely used concentration term is an amount per unit volume of fluid; that is, a volume-based concentration. But volume-based concentration is not conserved in a variable-density system. The term "solute concentration," used in this report, will refer to the mass-based concentration or mass fraction. The conservation equation for the solute in the fluid phase can be written:

$$\frac{\partial(\epsilon\rho\omega)}{\partial t} = \nabla \cdot \epsilon \rho \underline{D}_{S} \nabla \omega + \nabla \cdot \epsilon \rho \underline{D}_{m} \underline{I} \nabla \omega - \nabla \cdot \epsilon \rho \underline{v} \omega - \lambda \epsilon \rho \omega$$

$$-\rho_{b} R_{fs} + q \rho^{*} \omega^{*}; \qquad (2.1.3.1a)$$

\* where

w is the mass fraction of solute in the fluid phase (-);
w\* is the mass fraction of solute in the fluid source (-);
<u>P</u><sub>S</sub> is the mechanical-dispersion-coefficient tensor (m<sup>2</sup>/s);
D is the effective-molecular diffusivity of the solute (m<sup>2</sup>/s);
λ is the linear-decay rate constant (s<sup>-1</sup>);
R<sub>fs</sub> is the transfer rate of solute from fluid to solid phase per unit mass of solid phase (kg solute/s·kg solid phase); and
ρ is the bulk density of the porous medium (kg/m<sup>3</sup>).

A similar conservation equation can be written for the solute in the solid phase:

$$\frac{\partial(\rho_b \bar{w})}{\partial t} = \rho_b R_{fs} - \lambda \rho_b \bar{w} , \qquad (2.1.3.1b)$$

where

 $\bar{w}$  is the mass fraction of solute on the solid phase (-).

The solute is immobile when it is on the solid phase. Under the assumption of linear-equilibrium sorption, the fluid-phase and solid-phase concentrations can be related by an equilibrium-distribution coefficient:

 $\vec{w} = K_{d} \rho w$ ; (2.1.3.1c)

where

 $K_d$  is the equilibrium-distribution coefficient (m<sup>3</sup>/kg).

By combining equations 2.1.3.1a-c, we obtain the final soluteconservation equation:

$$\frac{\partial}{\partial t} (\varepsilon + \rho_{b}K_{d})\rho w = \nabla \cdot \varepsilon \rho [\underline{\underline{D}}_{S} + \underline{D}_{m}\underline{\underline{I}}] \nabla w - \nabla \cdot \varepsilon \rho \underline{v} w - \lambda (\varepsilon + \rho_{b}K_{d})\rho w$$

+qρ\*w\* ;

(2.1.3.2)

Equation 2.1.3.2 relates the rate-of-change of solute in the fluid phase to the net dispersive and diffusive flux, the net advective flux, the solutesource rate, the solute-injection rate with a fluid source, and the solutedecay rate. The equation is written for a unit volume of fluid and solid phase together; that is, a unit volume of saturated porous medium. Note that solute is injected into the sytem at concentration, w\*, and density,  $\rho$ \*, by a fluid source; but that solute is withdrawn at concentration w, and density  $\rho$ , by a fluid sink; that is, w\* = w, if q<0.

## 2.2. PROPERTY FUNCTIONS AND TRANSPORT COEFFICIENTS

Before the three conservation equations can be solved, information about the fluid properties, porous-matrix properties, and transport coefficients need to be obtained. The fluid properties are density, viscosity, heat capacity, thermal conductivity, and reference-state enthalpy. The porousmatrix properties are porosity, compressibility, permeability, heat capacity, thermal conductivity, and reference-state enthalpy. The transport coefficients are heat- and solute-dispersion tensors, and the effective molecular diffusivity, decay and sorption coefficients of the solute. In the HST3D simulator, density, viscosity, and porosity are functions of the dependent variables: pressure, temperature, and solute-mass fraction. The heat- and solute-dispersion tensors are functions of space and the interstitial velocity. The other parameters are either uniform or functions of space within the simulation region.

## 2.2.1. Fluid-Density Function

Fluid density is assumed to be a function of pressure, temperature, and solute concentration. For fluids such as water, a linear-density function is usually adequate over the ranges of pressures, temperatures, and solute concentrations encountered. Thus, the fluid-density function incorporated into this simulation code is:

$$\rho(\mathbf{p},\mathbf{T},\mathbf{w}) = \rho(\mathbf{p}_{o},\mathbf{T}_{o},\mathbf{w}_{o}) + \frac{\partial\rho}{\partial\mu} \Big|_{o}(\mathbf{p}-\mathbf{p}_{o}) + \frac{\partial\rho}{\partial\mathbf{T}} \Big|_{o}(\mathbf{T}-\mathbf{T}_{o}) + \frac{\partial\rho}{\partial\omega} \Big|_{o}(\mathbf{w}-\mathbf{w}_{o}); \qquad (2.2.1.1a)$$

or

2

$$\rho(p,T,w) = \rho_{o} + \rho_{o}\beta_{p}(p-p_{o}) - \rho_{o}\beta_{T}(T-T_{o}) + \rho_{o}\beta_{w}(w-w_{o}) ; \quad (2.2.1.1b)$$

where

ρ<sub>o</sub> is the fluid density at a reference pressure, p<sub>o</sub>, temperature, T<sub>o</sub>, and mass fraction, w<sub>o</sub>, (kg/m<sup>3</sup>);
 β<sub>p</sub> is the fluid compressibility (Pa<sup>-1</sup>);
 β<sub>T</sub> is the fluid coefficient of thermal expansion (°C<sup>-1</sup>); and
 β<sub>w</sub> is the slope of the fluid density as a function of mass fraction divided by the reference fluid density (-).

Now  $\rho_{\alpha}\beta_{\omega}$  is given by:

$$\rho_{o}\beta_{w}^{c} = \frac{\rho(w_{max}) - \rho(w_{min})}{w_{max} - w_{min}} | p_{o}, T_{o}$$
(2.2.1.1c)

where

The user needs to specify  $w_{\min}$  and  $w_{\max}$  along with  $\rho(w_{\min})$  and  $\rho(w_{\max})$ . The minimum solute-mass fraction usually will be determined by the initial conditions. If linear decay is present,  $w_{\min}$  must be zero. The maximum solute-mass fraction usually will be determined by source or boundary conditions because none of the transport processes incorporated in the HST3D simulator will concentrate solute in the fluid phase. For simplicity,  $w_0$  is taken to be equal to  $w_{\min}$ .

The option is available in HST3D to use a scaled, solute, mass fraction defined by:

$$w' = \frac{w - w_{\min}}{w_{\max} - w_{\min}}; \qquad (2.2.1.2)$$

where

w' is the scaled solute-mass fraction (-);

The scaled solute-mass fraction also is dimensionless and ranges from 0 to 1. Commonly, for input and output of mass-fraction data, it is more convenient to deal with a scaled solute-mass fraction rather than an absolute value. With a scaled solute-mass fraction, equation 2.2.1.1b becomes:

$$\rho(p,T,w') = \rho_{o} + \rho_{o}\beta_{p}(p-p_{o}) - \rho_{o}\beta_{T}(T-T_{o}) + \rho_{o}\beta_{w}'w', \quad (2.2.1.3a)$$

where

$$\rho_{o}\beta_{w}' = \rho(w_{max}) - \rho(w_{min})$$
 (2.2.1.3b)

The errors caused by assuming constant values for fluid compressibility, coefficient of thermal expansion, and variation of density with solute concentration can be assessed by looking at a density table for salt brines (Perry and others, 1963, p. 3-77).

Over a temperature range of about 100 °C and a solute-mass fraction range of 20 percent, the coefficient of thermal expansion varies by 60 percent and the density-concentration coefficient,  $\beta_w$ , varies by about 10 percent (Perry and others, 1963, p. 3-77). The variation of the fluid compressibility could not be checked because of lack of data. However, the density dependence on pressure for nearly incompressible fluids like water is much less than the density dependence on temperature or solute concentration. Therefore, some error will be introduced into the simulations by the linear-density function where large variations in temperature and solute concentration are involved.

The relative importance of pressure, temperature, and solute concentration for density variation can be seen from the salt-brine density table given in Perry and others (1963) and the compressibility of water. A change in pressure of  $10^6$  Pa results in a density change of about 0.04 percent, whereas a change in temperature of 100 °C results in a density change of about 4 percent, but a change in solute-mass fraction of 0.25 results a density change of about 20 percent. Thus, the salt concentration has the greatest effect on the density for typical ranges of the variables.

#### 2.2.2. Fluid-Viscosity Function

Fluid viscosity is strongly dependent on temperature, and, to a lesser extent, on solute concentration. The viscosity dependence on pressure is neglected. The viscosity as a function of temperature and scaled-solute concentration is written as:

$$\mu(T,w') = 10^{-3} \mu(T_{ov},w') \exp\left[(B_{0}w' + B_{1}(1-w'))(\frac{1}{T} - \frac{1}{T_{ov}})\right], (2.2.2.1)$$

where

µ(T<sub>ov</sub>, w') is the fluid viscosity at the reference temperature (kg/m-s); B<sub>0</sub>, B<sub>1</sub> are parameters describing the temperature dependence of viscosity at the concentration extremes (°C); and T<sub>ov</sub> is the reference temperature for viscosity (°C). The scaled solute-mass fraction of equation 2.2.1.2 is used in the viscosity function as well as the density function. The parameters  $B_0$  and  $B_1$  are obtained from a least-squares fit of viscosity versus temperature data. If jata are available only at a single temperature, the generalized viscosity versus temperature graph of Lewis and Squire as given in Perry and others (1963, p. 3-228) is used.

The concentration extremes are chosen to be the same minimum and maximum mass fractions described in section 2.2.1. The variation of viscosity with solute-mass fraction is specified in tabular form by the user. If viscosity data at only the minimum and maximum mass-fraction values are available, the equation used for viscosity as a function of concentration at a given temperature is:

$$\mu(w') = \mu_1(T_{ov})^{w'} \mu_0(T_{ov})^{1-w'}, \qquad (2.2.2.2)$$

where

- $\mu_0$  is the viscosity at the minimum-mass fraction or scaled concentration of zero (kg/m-s); and
  - $\mu_1$  is the viscosity of the maximum-mass fraction or scaled concentration of one (kg/m-s).

Equation 2.2.2.2 is used with equation 2.2.2.1 or alone in the case of isothermal simulation.

The viscosity versus temperature and concentration data that could be available may be divided into three classes. Class 1 is the greatest amount available, namely  $\mu(T)$  at  $w_{min}$  and  $w_{max}$  and  $\mu(w)$  for a range of w from  $w_{min}$  to  $w_{max}$ . Class 2 is viscosity versus temperature,  $\mu(T)$ , at only  $w_{min}$  and  $w_{max}$ . Class 3 is the least amount of data required, namely two viscosity points at a given temperature at  $w_{min}$  and  $w_{max}$ .

An evaluation of the accuracy of viscosity functions given in equations 2.2.2.1 and 2.2.2.2 was presented by INTERCOMP Resource Development and

Engineering, Inc. (1976). They found errors ranging from 5 to 14 percent over the temperature range from freezing to beiling for pure water. For a solution of sodium chloride with a mass traction ranging from 0.0 to 0.24, the different amounts of data available resulted in errors from 5 to 18 percent at a temperature of 65 °C. A sucrose solution with mass fractions ranging from 0.0 to 0.5 showed a maximum viscosity error of 30 percent. Other viscosity functions of temperature and solute concentration may be more suitable for certain situations.

## 2.2.3. Fluid Enthalpy

Fluid-phase enthalpy is a function of pressure, temperature, and solute concentration. The present version of the HST3D code uses the enthalpy of pure water obtained from the steam tables of Keenan and others (1969, p. 2-7 and 104-107), which can be described as:

$$H(\hat{p},T) = H(\hat{p}_{sat},0) + \int_{\hat{p}_{sat}}^{\hat{p}} \frac{1}{\rho} [1-\hat{T}\beta_{T}]d\hat{p} + \int_{o}^{T} c_{fo}dT; \qquad (2.2.3.1a)$$

where

H is the specific enthalpy of the fluid phase (J/kg); p̂ is the absolute pressure (Pa); p̂<sub>sat</sub> is the absolute pressure at saturation (Pa); and T̂ is the absolute temperature (K). c<sub>fo</sub> is the heat capacity of pure water at constant pressure (J/kg-°C).

The reference state for the enthalpy tables is saturated liquid water at 0 °C where the reference enthalpy is taken to be zero (Van Wylen, 1959, p. 80). The variation of enthalpy with solute concentration is treated in an approximate fashion, by adjusting the pure-water enthalpy by a factor that is the ratio of the heat capacity of the solution to the heat capacity of pure water at 0 °C, and by using an average heat capacity for the range of solute concentrations to be simulated. The heat capacity is assumed independent of temperature and pressure.

Thus,

$$H(p,T,w) = H(p,T,0) \ (\overline{c_f(w)}/c_{fo})$$

where

 $\overline{c_{\epsilon}(w)}$  is an average heat capacity (J/kg-°C).

During the simulations, the enthalpy is calculated as a variation from a reference state described by a pressure,  $P_{oH}$ , and a temperature,  $T_{oH}$ , selected by the user. The reference state is pure water so the reference mass fraction,  $w_{oH}$ , is always zero. Thus, the enthalpy equation becomes:

$$H(p,T,w) = H(\hat{p}_{oH}, T_{oH}, 0) \ (\overline{c_f}/c_{fo}) + \int_{p_{oH}}^{p} [1-\hat{T}\beta_T] \ \frac{dp}{\rho} + \int_{T_{oH}}^{T} \overline{c_f} dT \ ; \ (2.2.3.1c)$$

where

 $p_{oH}$  is a reference pressure for enthalpy (Pa);  $\hat{p}_{oH}$  is the corresponding absolute pressure (Pa); and  $T_{oH}$  is a reference temperature for enthalpy (°C).

The  $\hat{T}\beta_{T}$  term may be neglected for temperatures less than 100 °C (373 K) and density may be regarded as constant for pressure changes less than 10<sup>8</sup> Pa. The chosen reference pressure and temperature needs to be within the range to be calculated during the simulation. The heat capacity of the fluid needs to be an average value over the solute-concentration range to be simulated. More sophisticated treatments of the enthalpy of fluid mixtures are available in the literature; for example, Hougen and others (1959, p. 879).

#### 2.2.4. Porous-Medium Enthalpy

Enthalpy of the porous medium is taken to be a function of only temperature in the following form:

$$H_{s} = H_{s}(T_{oH}) + c_{s}(T - T_{oH}) ; \qquad (2.2.4.1)$$

where

H is the specific enthalpy of the solid phase (porous matrix) (J/kg); and

c\_ is the heat capacity of the solid phase (porous matrix)  $(J/kg^{\circ}C)$ .

Often, the enthalpy of the porous matrix is taken to be zero at a reference state of 0  $^{\circ}$ C.

### 2.2.5. Porous-Medium Compressibility

Many types of compressibility for porous media have been defined (Bear, 1972, p. 52, 203-213; Thomas, 1982, p. 34, 40). The porous-medium bulk compressibility,  $\alpha_{\rm b}$  (Pa<sup>-1</sup>), is defined on a volumetric basis (Bear, 1972, p. 56; Eagleson, 1970, p. 268), assuming confined-aquifer conditions, and one-dimensional, vertical consolidation of the porous matrix, as:

$$\alpha_{b} = \frac{1}{V_{b}\partial p}^{\partial V_{b}}; \qquad (2.2.5.1)$$

where

 $V_b$  is the bulk or total volume of a fixed mass of porous medium, that is, fluid plus porous matrix (m<sup>3</sup>).

Petroleum-reservoir engineers use the term rock compressibility,  $\alpha_r$  (Pa<sup>-1</sup>), defined as (Thomas, 1982, p. 34):

$$\alpha_{\rm r} = \frac{1}{\varepsilon} \frac{\partial \varepsilon}{\partial p} \qquad (2.2.5.2)$$

Rock compressibility directly expresses the variation of porosity with pressure. It is related to bulk compressibility by:

$$\alpha_{\rm r} = \frac{(1-\varepsilon)}{\varepsilon} \alpha_{\rm b} ; \qquad (2.2.5.3a)$$

> for the case of a nondeforming control volume, where more porous medium enters the control volume, as compression takes place. It is related by:

$$\alpha_r = \frac{\alpha_b}{\varepsilon}$$
(2:2.5.3b)

for the case of a deforming control volume, or where impermeable medium enters a nondeforming control volume, as compression takes place.

By combining equations 2.2.5.2 and 2.2.5.3b we obtain:

$$\frac{\partial \varepsilon}{\partial p} = \alpha ; \qquad (2.2.5.4)$$

which relates bulk compressibility to changes in porosity with changes in pressure.

Thus we have allowed the control volume to deform as the porous matrix and the fluid specific volumes expand or contract with changes in pressure. However, we neglect the velocity of deformation, so that the interstitial-pore velocity is calculated with respect to the fixed-coordinate system.

The specific storage is related to the compressibilities of the fluid and porous medium by (Eagleson, 1970, P. 270):

 $S_{o} = \rho g(\alpha_{b} + \epsilon \beta_{p})$ 

(2.2.5.5)

where

•

S is the specific storage  $(m^{-1})$ .

However, it is more convenient for our purposes to employ the conpressibility parameters, because of the variable density.

2.2.6. Dispersion Coefficients

## 2.2.6.1. Solute Dispersion

Hydrodynamic dispersion is the name for the group of mixing mechanisms that occur on the micro or pore scale that cause the irreversible spreading of a solute tracer that is observed at the macro or field scale for the system. As described by Bear (1972, p. 580-581), flow within the porous-medium structure has variations in local flow velocity, because of the velocity profile across the pore and mixing at pore junctions. The macroscopic effect is mechanical dispersion of a tracer. Molecular diffusion also is present where solute-tracer concentration gradients exist. However, diffusion in liquids is a relatively slow process, producing significant transport rates only at very slow ground-water flow velocities. In a laminar flow regime within the pores, diffusion of solute from one flow path to another contributes to the dispersion, so the separation of dispersion into a mechanical and diffusive mechanisms is somewhat artificial. For an extensive discussion of dispersion theory and a review of previous work, see Bear (1971, ch. 10).

The form of the hydrodynamic-dispersion-coefficient tensor  $D_{Sij}^{T}$  (m<sup>2</sup>/s) for the heat- and solute-transport simulation model is assumed to be, in component form:

$$D_{sij}^{*} = D_{sij}^{*} + D_{m}^{\delta}_{ij};$$
 (2.2.6.1.1)

where

 $D_{sij}$  is the mechanical-dispersion-tensor component  $(m^2/s)$ ;  $D_m$  is the effective molecular-diffusion coefficient  $(m^2/s)$ ; and  $\delta_{ij}$  is the Kronecker delta function.

The effective molecular-diffusion coefficient is the liquid-phase molecular diffusivity multiplied by an attenuation factor that accounts for the effect of the tortuosity of the porous medium. The form of the mechanical-dispersion coefficient is taken from the work of Scheidegger (1961) and Bear (1961) as presented by Konikow and Grove (1977) and Bear (1972, ch. 10). For an isotropic porous medium, two parameters describe the mechanical-dispersion tensor, the longitudinal dispersivity,  $\alpha_{T}$  (m), and the transverse dispersivity,  $\alpha_{T}$  (m). Then the nine components of the mechanicaldispersion tensor are given by:

$$D_{\text{Sij}} = (\alpha_{\text{L}} - \alpha_{\text{T}}) \frac{\frac{v_{i}v_{j}}{v}}{v} + \alpha_{\text{T}} v\delta_{ij}; \qquad (2.2.6.1.2)$$

(2.2.6.1.3)

where

v, is the component of interstitial velocity in the ith direction . (m/s):

and

where

v is the magnitude of the velocity vector (m/s).

 $v = (v_1^2 + v_2^2 + v_3^2);$ 

In general, the subscript 1 is associated with the x direction; the subscript 2 is associated with the y direction; and the subscript 3 is associated with the z direction. Field data have shown that longitudinal dispersivity usually is 3 to 10 times larger than transverse dispersivity (Freeze and Cherry, 1979, p. 396; Anderson, 1979), and that their magnitudes are dependent on the scale of observation distance over which the tracer is transported in the system.

Note that while flow in the porous medium may be governed by an anisotropic-permeability tensor, dispersion for heat and solute transport is assumed to be described by a dispersion tensor that applies to an isotropicporous medium. This assumption is made because it is not feasible to obtain all the dispersivity parameters for an anisotropic medium. If dispersive transport is a second-order effect, relative to advective transport, this inconsistency should not introduce serious errors. In most cases, the errors should be less than those introduced by uncertainties in the dispersion parameters themselves.

When the longitudinal and transverse dispersivities are not equal, dispersive transport will cause a solute distribution to enlongate in the direction of flow, because the longitudinal dispersivity always is greater than or equal to the transverse dispersivity. Thus, anisotropic spreading of solute and heat can occur in an isotropic-porous medium, even under conditions of uniform, unidirectional flow.

### 2.2.6.2. Thermal Dispersion

A description of thermal dispersion is based on a direct analogy with solute dispersion. Energy replaces solute mass as the quantity being transported by mechanical dispersion, and thermal conduction replaces molecular diffusion. Thus, the thermo-mechanical dispersion tensor is derived from the mechanical dispersion tensor by:

$$D_{Hij} = \rho c_f D_{Sij};$$
 (2.2.6.2.1)

where

 $D_{Hij}$  is the thermo-mechanical-dispersion tensor component (W/m-°C).

Combining the thermo-mechanical dispersion tensor with the net thermal conductivity of the fluid and solid phases gives the thermo-hydrodynamicdispersion coefficient tensor,  $D_{Hij}^{*}$  (W/m-°C), in component form:

$$D_{Hij}^{*} = D_{Hij} + [\varepsilon K_{f} + (1-\varepsilon)K_{s}]\delta_{ij}$$

(2.2.6.2.2)

# 2.3. EXPANDED SYSTEM EQUATIONS

When the density function, equation 2.2.1.1b, and the porous-medium compressibility relation, equations 2.2.5.3a and 2.2.5.3b are incorporated into the system governing equations, the following expanded system equations are obtained:

For ground-water flow:

$$\epsilon \rho_0 \beta_p \frac{\partial p}{\partial t} + \epsilon \rho_0 \beta_T \frac{\partial T}{\partial t} + \epsilon \rho_0 \beta_w \frac{\partial w}{\partial t} + \rho \alpha_L \frac{\partial p}{\partial t} = \nabla \cdot \rho \frac{k}{w} (\nabla p + \rho g) + q \rho^* ; \quad (2.3.1s) -$$

For heat transport:

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$$\epsilon \rho_{o} \beta_{p} c_{f} T \frac{\partial p}{\partial t} + \epsilon \rho_{o} \beta_{T} c_{f} T \frac{\partial T}{\partial t}$$

$$+ \epsilon \rho_{o} \beta_{w} c_{f} T \frac{\partial w}{\partial t} + \rho \alpha_{b} c_{f} T \frac{\partial p}{\partial t}$$

$$+ \epsilon \rho c_{f} \frac{\partial T}{\partial t} - \rho_{s} c_{s} T \alpha_{b} \frac{\partial p}{\partial t} + (1 - \epsilon) \rho_{s} c_{s} \frac{\partial T}{\partial t}$$

 $= \nabla \cdot (\varepsilon K_{f} + (1-\varepsilon)K_{s}) I \nabla T$ 

+ 
$$\nabla \cdot \varepsilon \underline{D}_{u} \nabla T - \nabla \cdot \varepsilon \rho c_{f} v T$$

+  $q_{\rm H}$  +  $q_{\rm p} * c_{\rm f} T^*$ ; (2.3.1b)

For solute transport:

$$\rho_{o}\beta_{p}(\epsilon+\rho_{b}K_{d}) = \frac{\partial p}{\partial t} + \rho_{o}\beta_{T}(\epsilon+\rho_{b}K_{d}) = \frac{\partial T}{\partial t}$$

$$+ \rho_{o}\beta_{w}(\epsilon+\rho_{b}K_{d}) = \frac{\partial w}{\partial t}$$

$$+ \rho \alpha_{b} = \frac{\partial p}{\partial t} + \rho(\epsilon+\rho_{b}K_{d}) = \nabla \cdot \epsilon \rho [\underline{p}_{S} + D_{m}\underline{I}] \nabla w - \nabla \cdot \epsilon \rho \underline{v} w$$

$$- \lambda(\epsilon+\rho_{b}K_{d}) \rho w + q \rho^{*} w^{*} \qquad (2.3.1c)$$

The change in the product of bulk density and equilibrium-distribution coefficient,  $\rho_b K_d$ , with pressure is 2 ro, because these equations were derived for a fixed mass of porous medium occupying a volume that under-goes slight deformation with variations in pressure. These three expanded equations show \_ the implicit coupling that occurs with variable density and porosity.

### 2.4. SOURCE OR SINK TERMS--THE WELL MODEL

Most of the ground-water flow and heat and solute sources or sinks affect the simulations through the boundary conditions. However, a line source or sink term is used to represent injection or withdrawal by a well. Although a well is treated as a line source or sink for the flow and transport equations, a well is a finite-radius cylinder for the well-bore model.

The well model for the HST3D simulator is more sophisticated than those well models used in most ground-water flow simulators. A well can be used for fluid injection or fluid withdrawal, with associated heat and solute injection or production. It also can be used simply for observation of aquifer conditions. In the present code, the well bore can communicate with any subset of cells along the z-coordinate direction at a given x-y location. That is, the well may be screened or it may be an open hole over several intervals of its depth. Several options are available for specifying pressure

or flow-rate conditions under which the well will operate. A special technique is used to relate the local pressure field around a well to the pressures in the cells with which it communicates. Finally, a mathematical model of the well riser is included to calculate pressure and heat gains and losses as fluid moves from the land surface to the uppermost screened interval, or vice versa.

The well can be divided into two parts as shown in figure 2.1. The lower part, from the bottom of the borehole to the top of the uppermost screened interval, will be referred to as the well bore; the upper part, from the top of the screened interval to the land surface, will be referred to as the well riser. The well-riser interval may or may not have a riser pipe within it, and the well-bore interval may be an open hole or have cased and screened sections. A screened section also may be just perforated casing. The term well-datum level refers to the location at the junction between the well riser and the well bore, equivalently referred to as the bottom hole.

Focusing attention on the well bore, we shall describe the linking of the well model to the simulation region as a source or sink, and then describe the pressure and flow-rate conditions that can be specified as bottom-hole conditions. The incorporation of the well-riser calculations will then be discussed.

Cell or nodal pressures represent a spatially averaged condition, when the simulation region is discretized into finite-difference cells. A well located in a cell will have a pressure at the screen at the nodal elevation that is not necessarily the same as the cell pressure. Various analytical approaches have been used to avoid the computational burden of a finer finite-difference grid around each well in the region. They are summarized by Aziz and Settari (1979, sec. 7.7) and are based on steady-state radial flow in a cylindrical-coordinate system with homogeneous aquifer properties. Another review may be found in Williamson and Chappelear (1981).

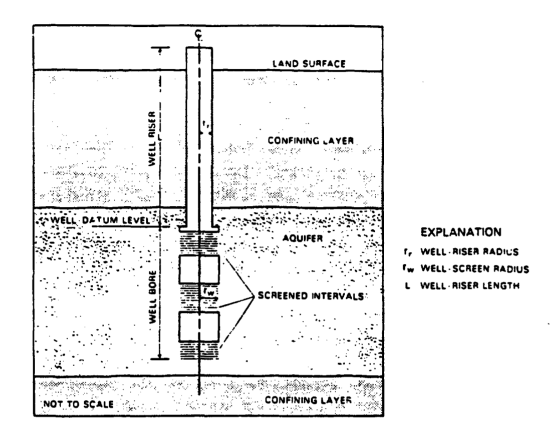


Figure 2.1.--Sketch of well-model geometry showing the well-bore and well-riser sections and the well-datum level.

## 2.4.1. The Well-Bore Model

For three-dimensional cartesian coordinates, the present version of the HST3D code uses a modification of the well-bore equation derived by Van Poolen and others (1968). Consider steady-state radial flow from a well into a homogeneous aquifer with flux across an exterior cylindrical boundary,  $r_e$ . This boundary can be regarded as a radius of influence of the well. For a cartesian-coordinate system, the exterior radius,  $r_e$ , is taken to be the

radius of a circle that encloses the equivalent area to the x-y horizontal area of the cell in which the well is located. The average pressure within the annulus between the well-bore radius and the radius of influence can be calculated and the flow rate from the well per unit length of well bore can be expressed as a function of the pressure change from the well-bore pressure to this average pressure. At any given elevation, z, we have:

$$q_{w} = \frac{2\pi k_{w} (r_{e}^{2} - r_{w}^{2}) (p_{w} - p_{av})}{\mu [r_{e}^{2} \ln (r_{e}/r_{w}) - 0.5 (r_{e}^{2} - r_{w}^{2})]}; \qquad (2.4.1.1)$$

where

p<sub>w</sub> is the pressure at the wel! bore (Pa);
k<sub>w</sub> is the average permeability between r<sub>w</sub> and r<sub>e</sub> (m<sup>2</sup>);
p<sub>av</sub> is the average pressure between r<sub>w</sub> and r<sub>e</sub> (Pa);
r<sub>w</sub> is the well-bore radius (m);
r<sub>e</sub> is the radius of influence of the well (m); and
q<sub>w</sub> is the volumetric flow rate per unit length of well bore
 (positive is flow into the aquifer) (m<sup>3</sup>/m-s).

The time-independent factors that affect flow from a well bore can be combined into a single term. Departing slightly from petroleum-reservoirengineering usage, we define a modified well index as follows:

$$W_{I} = \frac{2\pi k_{w} (r_{e}^{2} - r_{w}^{2})}{r_{e}^{2} \ln (r_{e}/r_{w}) - 0.5 (r_{e}^{2} - r_{w}^{2})}; \qquad (2.4.1.2)$$

where

 $W_{\tau}$  is the well index per unit length of well bore (m<sup>2</sup>).

The average permeability k is taken to be:

$$k_w = (k_x k_y)^{\frac{1}{2}};$$
 (2.4.1.3)

for cartesian-coordinate systems, where

 $k_{\chi}$  is the permeability in the x-direction (m<sup>2</sup>); and  $k_{\chi}$  is the permeability in the y-direction (m<sup>2</sup>).

There is presently no provision for accommodating areally heterogeneous permeability distributions in the vicinity of the well bore.

Equations 2.4.1.1 to 2.4.1.3 will be modified for use with the finitedifference discretization in the numerical-implementation section 3.3.

For three-dimensional and cylindrical regions, the total specified flow rate from the well needs to be allocated over the length of well bore that communicates with the aquifer. This allocation can be done in two ways; by fluid mobility, or by the product of fluid mobility and the pressure difference between the aquifer and the well bore. Although there may be zones of cased well bore through which there is no communication with the aquifer, we shall assume for the present discussion that the well bore is screened throughout its depth. The total well flow rate from the well to the aquifer is given by:

 $Q_{u} = \int_{L}^{L} q_{u} dl;$  (2.4.1.4a)

$$= \int_{\frac{Q}{L}}^{\frac{Q}{L}} \frac{w_{I}(2)}{\mu(2)} (p_{w} - p_{av}) d2 ; \qquad (2.4.1.4b)$$

Q<sub>w</sub> is the volumetric well flow rate (positive is from the well to the aquifer) (m<sup>3</sup>/s);

- L is the distance along the well bore (m);
- $\boldsymbol{\ell}_{T}$  is the lower end of the screened interval (m); and
- $\ell_{11}$  is the upper end of the screened interval (m).

Fluid mobility at the well can be defined as:

$$M_{w}(l) = \frac{W_{I}(l)}{\mu(l)};$$
 (2.4.1.5)

where

M<sub>1</sub> is the well mobility per unit length of well bore  $(m^3/s-m-Pa)$ .

Allocation of the specified flow rate by fluid mobility is obtained by assuming that the pressure difference in equation 2.4.1.4b is independent of depth. Then

$$q_{w}(\ell) = M_{w}(\ell) / \int_{\ell_{T}}^{\ell_{U}} M_{w}(\ell) d\ell ;$$
 (2.4.1.6)

represents the allocation of the total flow rate over the well-bore length as a function of fluid mobility.

For wells drilled at an angle,  $\theta_{ij}$ , to the vertical or z-axis,

 $dz = \cos \theta_{ij} dl;$ 

(2.4.1.7)

 $v_{ij}$  is the angle between the vertical and the well bore (degrees).

If the screened interval is not continuous from  $\ell_L$  to  $\ell_U$ , the mobility is set to zero over the appropriate subintervals.

The alternative method of flow-rate allocation over the well-bore length is derived by not regarding the pressure difference in equation 2.4.1.4b as constant with depth. A hydrostatic-pressure distribution in the well bore is assumed using an average fluid density. Thus, frictional hydraulic-head losses in the well bore are neglected. This yields, from equation 2.4.1.4b.

$$P_{wd} = \frac{\int_{\ell_L}^{\ell_U} M_w(\ell) [p_{av}(\ell) + \rho_w g(z - z_{wd})] d\ell + Q_w}{\int_{\ell_L}^{\ell_U} M_w(\ell) d\ell}; \qquad (2.4.1.8)$$

where

 $p_{wd}$  is the bottom-hole or well-datum pressure (Pa);  $z_{wd}$  is the elevation of the well datum (m); and  $\rho_w$  is the average fluid density in the well bore (kg/m<sup>3</sup>).

Then the well flow rate is allocated as follows:

$$q_{w}(l) = M_{w}(l) [p_{wd} + \rho_{w}g(z_{wd} - z) - p_{av}];$$
 (2.4.1.9)

This method is referred to as allocation by mobility and pressure difference. The average pressure,  $p_{av}$ , will be related to the grid-cell pressures in section 3.3.1 on numerical implementation.

The flow rate can be specified with a bottom-hole pressure-constraint condition, that may affect the source or sink flow rate applied. Allocation is by mobility and pressure difference, and equation 2.4.1.8 is used to calculate a predicted bottom-hole pressure based on the specified flow-rate. For an injection well, if the predicted pressure is greater than the bottomhole constraint pressure, then the well is pressure-limited, and the flow rate will be less than that specified. The flow rate will be reduced to meet the pressure constraint. If the predicted bottom-hole pressure is less than that specified, then the desired flow rate is used. For a production well, if the predicted bottom-hole pressure is less than the constraint pressure, the well is pressure limited, and the flow rate will be less than desired. Otherwise, the pressure constraint is not limiting. In other words, a well bore can function as either a Dirchlet or a Neumann boundary condition, or it can switch back and forth.

When bottom-hole (well-datum) pressure is specified, equation 2.4.1.9 gives the flow-rate allocation and equation 2.4.1.4b gives the total flow rate. No constraints are applied to the calculated flow rate.

After the flow rate has been established and allocated, heat-injection and solute-injection rates are determined from the bottom-hole pressure, specified-temperature, and specified solute-mass-fraction values. Heatwithdrawal and solute-withdrawal rates are determined by the ambient pressure, temperature, and solute-mass fraction in the aquifer for each cell that communicates with a well bore.

In the case of cylindrical coordinates with a single well at the radial origin, the inner radius of the simulation region becomes the well-bore surface. Thus, a specified flow rate allocated by mobility becomes a specified-flux boundary condition. Allocation by mobility and pressure difference using equation 2.4.1.9 is not applicable here, because the well-bore pressure and the pressure at the inner radius of the region are identical. Instead, the pressure profile along the well bore is not assumed to be hydrostatic, but, rather it satisfies a steady-state momentum equation, that includes frictional pressure losses, but neglects changes in momentum by flow into or out from the well bore. Then, we have, for a differentialmomentum balance along the well bore:

$$\frac{dp_{w}}{dz} + \rho_{w}g + \frac{\rho_{w}v^{2}_{w}}{4r_{w}} f_{w} = 0; \qquad (2.4.1.10)$$

f is the hydraulic-head-loss friction factor (-); and

v is the average velocity across the well bore at a given z-level (m/s).

The corresponding mass balance is obtained assuming no change in wellbore storage, thus:

$$\frac{d\rho v_{w}}{dz} + \frac{2\rho_{w} q_{Fw}}{r_{w}} = 0 ; \qquad (2.4.1.11)$$

where

 $q_{Fw}$  is the volumetric flux from the well bore  $(m^3/m^2-s)$ .

Equations 2.4.1.10 and 2.4.1.11 can be combined to give:

$$\rho_{w}q_{Fw} = \frac{d}{dz} \left[ \frac{2r_{w}^{2}}{v_{w}f_{w}} \left( \frac{dp_{w}}{dz} + \rho_{w}g \right) \right]; \qquad (2.4.1.12)$$

Equation 2.4.1.12 is combined with the flow equation 2.1.1.1a by assuming that the aquifer pressure and well-bore pressure are equal at the well-bore radius. The flow equation at the inner radius of the region becomes:

$$\frac{\partial \varepsilon \rho}{\partial t} = \nabla \cdot \rho \frac{k}{\mu} (\nabla p + \rho g) + \frac{\partial}{\partial z} \left[ \frac{2r_w^2}{v_w f_w} \left( \frac{\partial p}{\partial z} + \rho_w \varepsilon \right) \right]; \quad (2.4.1.13)$$

tor the parts of the inner radius that are screened. A fluid-flux boundary condition of zero applies over the cased-off intervals.

Thus, the flow equation is still in its original form, but the coefficients of pressure gradient in the z-direction, and of the gravity term, are augmented. The flow rate to or from the well is implicitly incorporated. When the equation is converted to discrete form, the flow rate to or from the well will arise naturally at the upper boundary of the screened interval. The friction-head-loss factor is calculated as described in the well-riser model, section 2.4.2. The magnitude of the friction head-loss factor often may be very small but it needs to be non-zero, for flow to occur in the well bore.

The total flow rate to or from the well always is satisfied by this calculation method, and the pressure at the top of the screened interval in the aquifer is identical to the well-datum-level pressure. Recall that these pressures are not necessarily equal in the line-source approach used with the cartesian coordinate system. An examination of the relative magnitudes of the terms for advective momentum and frictional head-loss in the full momentumbalance equation shows that, for a producing well with uniform inflow per unit length, the advective-momentum term dominates near the bottom of the screen. The frictional head-loss term dominates at distances above the bottom of the screen that are greater than about 1,000 times the well radius. Thus, a significant region exists in which both the momentum and frictional terms are of similar magnitude. However, a more rigorous development, retaining the momentum term, is beyond the scope of this work. The present development follows that of Aziz and Settari (1979, p. 337-341).

### 2.4.2. The Well-Riser Model

When flow rate or pressure is specified at the land surface for a given well, the well-riser calculation needs to be performed in conjunction with the well bore flow-rate allocation described above. This calculation consists of a simultaneous solution of the macroscopic equations of total energy, momentum and mass (Bird and others, 1960, p. 209-212) for the change in pressure and temperature over the well-riser length.

The total-energy or enthalpy equation is written for steady flow either up or down the well riser as a rate of change with distance along the riser,

$$\frac{dH_r}{d\ell} + g \cos\theta_r + v_r \frac{dv_r}{d\ell} = Q_{Hr}(\ell); \qquad (2.4.2.1)$$

where

H<sub>r</sub> is the specific enthalpy of fluid in the riser (J/kg);
ν<sub>r</sub> is the average velocity across the riser at a given *L*-location (m/s);
θ<sub>r</sub> is the angle between the well riser and vertical (degrees);
Q<sub>Hr</sub> is the heat transferred per unit mass per unit length to the fluid in the riser (J/kg-m); and *L* is the distance along the well-riser casing (m).

Energy loss by viscous dissipation has been neglected. All quantities are averages across the riser-pipe cross section at a given level.

The equation for momentum along the well-riser axis also is written for steady flow as a differential balance along the well riser:

$$2\rho_r v_r \frac{dv_r}{d\ell} + \rho_r g \cos\theta_r + \frac{dp_r}{d\ell} + \frac{\rho_r v_r^2}{2r_r} f_r = 0 ; \qquad (2.4.2.2)$$

 $\rho_{\_}$  is the fluid density in the ris r (kg/m³);

p\_ is the pressure in the riser (Pa);

 $r_{\mu}$  is the internal radius of the well riser (m); and

f\_ is the hydraulic-head-loss friction factor (-).

Finally, the macroscopic-mass balance, written in differential form as a rate of change along the riser, is:

$$\rho_r v_r = Q_{Fr} / \pi r_r^2;$$

(2.4.2.3a)

where

 $Q_{\rm Fr}$  is the total mass-flow rate in the riser (kg/s).

Differentiation with respect to length yields:

$$\rho_r \frac{dv_r}{d\ell} + v_r \frac{d\rho_r}{d\ell} = 0. \qquad (2.4.2.3b)$$

To solve equations 2.4.2.1, 2.4.2.2, 2.4.2.3a, and 2.4.2.3b, the enthalpy tables (Keenan and others, 1969, p. 2-7 and 104-107) are used for  $H_r(p,T)$ , equation 2.7a is used for the density equation of state, and the Fanning friction factor, using the Moody correlation (Perry and others, 1963, p.-5-20), is used to calculate  $f_r$  as a function of velocity. The enthalpy for pure water is adjusted for other fluid mixtures according to equation 2.2.3.1b. For turbulent flow, the friction factor is a function of pipe roughness. The user needs to supply a value for pipe roughness, and some typical values for pipe roughness from Shames (1962, p. 300) are given in table 2.1. Changes in viscosity with temperature along the riser are neglected.

Pipe type	Pipe roughness (millimeters)
Steel or wrought iron	$3.8 \times 10^{-3}$
Galvanized iron	$1.3 \times 10^{-2}$
Cast iron	$2.2 \times 10^{-2}$

Table 2.1.--Fipe-roughness values

The heat transferred to the fluid in the riser must pass from the surrounding medium to the riser pipe, then from the riser pipe to the fluid. The heat transferred per unit mass of fluid per unit length of riser is then:

$$Q_{Hr}(\ell) = \frac{2\pi r_r}{Q_{Fr}} U_T(T_a(\ell) - T_r(\ell));$$
 (2.4.2.4)

where

Tr is the fluid temperature in the well riser (°C);
Ta is the ambient temperature in the medium adjacent to the
riser (°C);
UT is the overall heat-transfer coefficient for the fluid, riser
pipe and surrounding medium (W/m<sup>2</sup>-°C).

The overall heat-transfer coefficient is given by:

$$\frac{1}{r_{r}U_{T}} = \frac{1}{r_{r}h_{r}} + \frac{\Delta r_{r}}{r_{r}K_{r}} + \frac{1}{K_{re}F_{CJ}(t)}; \qquad (2.4.2.5)$$

 $\Delta r_{p}$  is the wall thickness of the riser pipe (m);

 $F_{r,\tau}(t)$  is the dimensionless part of the Carslaw and Jaeger (1959,

- p. 336) solution for heat flux to an infinite medium from a constant-temperature cylindrical source (-);
- $h_r$  is the local heat-transfer coefficient from the fluid to the riser pipe (W/m<sup>2</sup>-°C);
- K is the thermal conductivity of the medium surrounding the riser pipe (W/m-°C); and
- K is the thermal conductivity of the riser pipe (W/m-°C).

Equation 2.4.2.5 is a simplification of the relation for the overall heat transfer coefficient for conduction through cylindrical walls (Bird and others, 1960, p. 288) combined with the Carslaw-Jaeger solution for heat flux to an infinite medium from a cylindrical source (Carslaw and Jaeger, 1959, p. 336). It is valid for wall thicknesses that are small relative to the riser-pipe radius.

The dimensionless heat-flux function,  $F_{CJ}(t)$ , can be approximated by the following two series:

(1) For short time, T, (Carslaw and Jaegar, 1959, p. 336):

$$F_{CJ} \cong F_{CJ}^{S}$$
; for  $\tau < 1$ ; (2.4.2.6a)

where

$$F_{CJ}^{S} \cong (\pi\tau)^{-\frac{1}{2}} + \frac{1}{2} - \frac{1}{2}(\frac{\tau}{\pi})^{\frac{1}{2}} + \frac{\tau}{8}; \qquad (2.4.2.6b)$$

and where

t is the dimensionless time defined by:

$$\tau = \frac{D_{\rm Hrm} t}{(r_{\rm r} + \Delta r_{\rm r})^2} ; \qquad (2.4.2.6c)$$

and where

 $D_{\text{Hrm}}$  is the thermal diffusivity of the medium surrounding the well riser  $(m^2/s)$ .

(2) For long times, the asymptotic expansion was derived by Ritchie and Sakakura (1956):

$$F_{CJ} \cong F_{CJ}^{L}$$
; for  $\tau > 3.6$ ; (2.4.2.7a)

where

$$F_{CJ}^{L} = 2(\ln \chi)^{-1} [1 - .5772(\ln \chi)^{-1} - 1.3118(\ln \chi)^{-2} + .2520 (\ln \chi)^{-3} + 3.9969 (\ln \chi)^{-4} + 5.0637 (\ln \chi)^{-5}] + \frac{4}{e^{2\gamma}} (\tau \ln \chi)^{-1} [(\ln \chi)^{-1} - 1.1544(\ln \chi)^{-2}] -2 \tau^{-1} (\ln \chi)^{-3} ; \qquad (2.4.2.7b)$$

$$\chi = \frac{4\tau}{e^{2\gamma}} ; \qquad (2.4.2.7c)$$

and where

 $\gamma$  is Euler's constant:  $\cong$  0.5772.

In equation 2.4.2.7b, terms of higher order than  $(\ln \chi)^{-6}$  and  $\tau^{-1}(\ln \chi)^{-3}$  have been dropped. Carslaw and Jaeger (1959, p. 336) present a lower-order version of equation 2.4.2.7b that is accurate for dimensionless time much greater than 3.6. The estimated error is on the order of 10 percent for dimensionless time, T, greater than 3.6. For a typical rock medium, this truncation means that the time must be greater than about 3.6  $\times$  10<sup>4</sup> s, or about 0.4 d. The short-time approximation, equation 2.4.2.6b, is good for time less than about 0.1 d. For intermediate time, the heat-transfer function is estimated by linear interpolation between  $F_{CJ}^{S}$  evaluated at T=1 and  $F_{CJ}^{L}$  evaluated at T=3.6.

Note that the heat-flux function in equation 2.4.2.5 is a function of. time; whereas, the mechanical and thermal-energy balances are at steady-state. This is a consistent approximation, provided it is assumed that the heat transfer from the fluid to the riser pipe and through to its outer boundary is rapid, relative to rates of change in temperature at the fluid-inlet end of the riser pipe; and, that changes in the fluid-temperature profile within the riser pipe re-equilibriate quickly, relative to induced temperature changes in the adjacent medium. This approach parallels that of Ramey (1962), with the difference being that the heat-flux solution from a cylinder at constant temperature is used, instead of the temperature solution for the constant heat-flux case. The former solution is considered to more accurately describe the physical situation.

Values for the local heat-transfer coefficient,  $h_r$  in equation 2.4.2.5, can be determined from correlations, such as those of McAdams (1954, p. 241-243) or Sieder and Tate (given in Bird and others, 1960, p. 399), between the Nusselt number, the Prandtl number, and the Reynolds number for forced convection in tubes.

The correlation from McAdams (1954, p. 219) that is valid for turbulent flow in the well-riser pipe is:

$$\frac{2r_{r}h_{r}}{\kappa_{f}} = 0.023 \left[ \frac{\rho_{r}v_{r}}{\mu_{r}} \right]^{0.8} \left[ \frac{c_{f}\mu_{r}}{\kappa_{f}} \right]^{0.33}; \qquad (2.4.2.8)$$

where

 $\mu_{\rm r}$  is the viscosity of the fluid in the riser pipe (kg/m-s).

The well-riser calculation is developed by combining equations 2.4.2.1-2.4.2.3b with equation 2.2.1.1b and the derivative of equation 2.2.3.1a for the enthalpy function. The resulting equations are:

$$\begin{bmatrix} v_{r}^{2} \beta_{p} - \frac{1}{\rho_{r}} & v_{r}^{2} \beta_{T} \\ & & & \\ & & & \\ \frac{\partial H}{\partial \rho}|_{T} - \frac{1}{\rho_{r}} & \frac{\partial H}{\partial T}|_{p} \end{bmatrix} \begin{bmatrix} \frac{dp_{r}}{dt} \\ & & \\ \frac{dT_{r}}{dt} \end{bmatrix} = \begin{bmatrix} g \cos \theta_{r} + \frac{v_{r}^{2}f_{r}}{2r_{r}} \\ & & \\ \frac{2\pi r_{r} U_{T}}{Q_{Fr}} & (T_{a} - T_{r}) + \frac{v_{r}^{2}f_{r}}{2r_{r}} \end{bmatrix} (2.4.2.9)$$

Using the thermodynamic relationships:

$$\frac{\partial H}{\partial p} \Big|_{T} = \frac{1}{\rho_{r}} - T \frac{\partial \rho_{r}^{-1}}{\partial T} \Big|_{p}; \qquad (2.4.2.10a)$$

and

 $\frac{\partial H}{\partial T}\Big|_{p} = c_{f}; \qquad (2.4.2.10b)$ 

we can reduce equation 2.4.2.9 to two simultaneous ordinary differential equations:

$$\begin{bmatrix} \frac{dp_{r}}{d\ell} \\ \frac{dT_{r}}{d\ell} \end{bmatrix} = \begin{bmatrix} c_{f} & -\beta_{T} v_{r}^{2} \\ -\beta_{T} T_{r} / \rho_{r} & \beta_{p} v_{r}^{2} - 1 / \rho_{r} \end{bmatrix}$$

$$\begin{bmatrix} g \cos \theta_{r} + v_{r}^{2} f_{r} / 2r_{r} \\ \frac{2\pi r_{r}}{Q_{Fr}} [\frac{1}{r_{r}h_{r}} + \frac{\Delta r_{r}}{r_{r}K_{r}} + \frac{1}{K_{re}F_{CJ}(t)}]^{-1} (T_{a} - T_{r}) + v_{r}^{2} f_{r} / 2r_{r} \end{bmatrix}$$

$$\begin{bmatrix} (\beta_{p} v_{r}^{2} - 1 / \rho_{r})c_{f} - T_{r} \beta_{T}^{2} v_{r}^{2} / \rho_{r} \end{bmatrix}^{-1} (2.4.2.11)$$

These equations are coupled through the density, velocity, and temperature terms. The boundary conditions are known at one end of the riser. For injection:

at 
$$z = z_{LS}$$
;  $p = p_{inj}$ ;  $T = T_{inj}$ ; (2.4.2.12a)

For withdrawal:

. .

at 
$$z = z_{wd}$$
;  $p = p_{wd}$ ;  $T = T_{wd}$ ; (2.4.2.12b)

where

 $z_{wd}$  is the elevation of the well datum (m); and  $z_{LS}$  is the elevation of the land surface (m).

Equations 2.4.2.6a-c and 2.4.2.7a-c are used to evaluate the heat-transfer function  $F_{CJ}$ .

The mass, enthalpy, and mechanical-energy-balance equations are solved either up or down along the well riser, depending on the direction of fluid flow, to obtain the pressure and temperature at the riser bottom for injection conditions, or at the riser top for production conditions. Co-pling this well-riser calculation to the well-bore model enables specified pressure, temperature, and solute concentration, or specified flow-rate conditions at the land surface, to be employed.

When the flow rate at the land surface is specified as an injection, the surface temperature and solute concentration also need to be specified. The well-riser calculation will give the necessary surface pressure to achieve the specified flow rate. If a production or withdrawal flow rate is specified, the surface pressure, temperature, and solute concentration are determined by the well-bore and well-riser calculations.

When the surface pressure is specified, the well-bore and well-riser calculations determine the flow rate, surface temperature, and solute concentration for a production well. Surface temperature and solute concentration also need to be specified in the case of an injection well. The ambient-temperature profile with depth along the well riser is specified by the user.

A flow rate and pressure constraint at the surface can be specified and the slower of the specified flow rate or the flow rate that results from the specified-pressure constraint will be applied to the aquifer and apportioned as described previously.

A well also can be used as an observation well. In this case, none of the well-bore or well-riser calculations are necessary. The purpose of an observation well is to record dependent variable data (pressure, temperature, solute-mass fraction) for plotting versus time at the conclusion of the simulation. The recorded data are the aquifer values at the well-datum level, which is at the top of the uppermost screened interval.

In summary, a well can be a production well, an injection well, or an observation well. The flow rate can be specified with or without a pressure constraint, or the pressure can be specified either at the land surface or at the well-datum level. For three-dimensional cartesian coordinates, the allocation of the flow to each layer can be determined by the relative mobility of the layer, or by the product of the mobility times the pressure difference. For cylindrical coordinates, the allocation is determined by the product of the mobility times pressure difference, with allowance for gravitational effects, because the well-bore equations are solved simultaneously with the ground-water flow equations for the region adjacent to the screened intervals. Application of the well-flow terms for each layer to the ground-water flow equation can be explicit or semi-implicit in time for three-dimensional cartesian coordinates; it is fully implicit for cylindrical coordinates.

## 2.5. BOUNDARY CONDITIONS

## 2.5.1. Specified Pressure, Temperature, and Solute-Mass Fraction

The first type of boundary condition, known as a Dirchlet boundary condition, is a specified pressure condition for the ground-water flow equation, a specified temperature condition for the energy-transport equation, and a specified-mass fraction for the solute-transport equation. These conditions can be specified independently as functions of location and they also can vary independently with time. Mathematically, we have:

$$p = p_{p}(x,t), \text{ for } x \text{ on } S_{t_{1}}^{1};$$
 (2.5.1.1a)

 $T = T_{B}(x,t)$ , for x on  $S_{T}^{1}$ ; and (2.5.1.1b)

 $w = w_{\rm B} ({\bf x}, t), \text{ for } {\bf x} \text{ on } S^1_{w};$  (2.5.1.1c)

where

 $p_B$  is the pressure at the specified boundary (Pa);  $T_B$  is the temperature at the specified boundary (°C);  $w_B$  is the mass fraction at the specified boundary (-);  $S_p^1$  is the part of the boundary with specified pressure;  $S_T^1$  is the part of the boundary with specified temperature; and  $S_w^1$  is the part of the boundary with specified mass fraction.

Care needs to be used in specifying the temperature and mass fraction at fluid-outflow boundaries, because, on boundary surfaces across which fluid flow occurs, the advective transport of heat and solute is assumed to dominate over any diffusive or dispersive transport. Thus, it is physically unrealistic to specify a temperature or solute concentration at an outflow boundary because the ambient fluid will determine the temperature, and solute concentration there.

# 2.5.2. Specified-Flux Boundary Conditions

The default boundary condition for the numerical model is no fluid, heat, or solute flux across the boundary surfaces. Normal fluxes of fluid, heat, and solute, known as Neumann boundary conditions, can be specified over parts of the boundary as functions of time and location. However, they cannot be

specified independently, because, on boundary surfaces where a specified fluid flux exists, the advective transport of heat and solute is assumed to dominate over any specified diffusive or dispersive flux of these quantities. This assumption means that, on fluid-inflow boundaries, the temperature and mass fraction of the inflowing fluid needs to be specified. These specifications determine the heat and solute fluxes. At fluid-outflow boundaries, the temperature and mass fraction are determined by the ambient fluid values in the region, thus giving the heat and solute fluxes. Therefore, it is not physically realistic to specify temperatures and mass fractions at outflow boundaries. On boundary surfaces where no fluid flux is given, heat and solute fluxes may be specified. Heat fluxes represent thermal conduction and solute fluxes represent solute diffusion.

For the reasons discussed in section 2.5.1, it also is not physically realistic to specify dispersive heat or solute fluxes across boundary surfaces that have specified pressures. However, total heat or solute fluxes may be specified for inflow boundaries. These fluxes are the advective fluxes approaching the boundary from outside the region, and they are equal to the advective plus the dispersive fluxes leaving the boundary and entering the region. For outflow boundaries, the boundary condition requires that the dispersive fluxes be zero. Thus, only advective flux of heat and solute occurs at outflow boundaries. Again, the advective transport of heat and solute is assumed to dominate over dispersive flux.

• Specified fluxes are expressed mathematically as:

$$q_{Fn} = (q_{Fx}^{B}, q_{Fy}^{B}, q_{Fz}^{B})$$
 for x on  $S_{p}^{2}$ ; (2.5.2.1a)

$$q_{Hn} = (q_{Hx}^B, q_{Hy}^B, q_{Hz}^B)$$
 for x on  $S_T^2$ ; (2.5.2.1b)

$$q_{Sn} = (q_{Sx}^{B}, q_{Sy}^{B}, q_{Sz}^{B})$$
 for x on  $S_{w}^{2}$ ; (2.5.2.1c)

- $q_{Fi}^{B}$  is the component of the fluid flux in the ith direction at the boundary  $(m^{3}/m^{2}-s)$ ;
- $q_{\text{Hi}}^{D}$  is the component of the heat flux in the ith direction at the boundary (W/m<sup>2</sup>);
- $q_{Si}^{B}$  is the component of the solute flux in the ith direction at the boundary (kg/m<sup>2</sup>-s);
- q<sub>Fn</sub> is the normal component to the boundary surface of the fluid-flux vector (kg/m<sup>2</sup>-s);
- $q_{Hn}$  is the normal component to the boundary surface of the heat-flux vector (W/m<sup>2</sup>);
- q<sub>Sn</sub> is the normal component to the boundary surface of the soluteflux vector (kg/m<sup>2</sup>-s); and
- $S_u^2$  are parts of the boundary with specified-fluid, heat, or solute fluxes respectively; u = p,T,w.

Note that the specified-fluid flux,  $q_F$ , is given as a volumetric flux. A fluid density also needs to be specified for the case of inflow to the region. The density in the region at the boundary is used for computation of mass outflow rates. Also note that flux is a vector quantity with components expressed relative to the coordinate system of the simulation region. Examples of physical-boundary conditions that can be represented as specified-flux boundaries include infiltration from precipitation, lateral boundaries where the pressure gradients can be estimated, and simple steady-state flow fields where recharge- and discharge-boundary flow rates are known.

2.5.3. Leakage Boundary Conditions

A leakage boundary condition has the property that a fluid flux occurs in response to a difference in pressure and gravitational potential across a confining layer of finite thickness. Usually the permeability of this layer will be orders of magnitude smaller than the permeability of the simulation region and the aquifer region on the other side of the confining layer.

Representation of leakage boundary conditions is based on the approach of Prickett and Lonnquist (1971, p. 30-35), which has been generalized to include variable-density and variable-viscosity flow. The mathematical treatment of leakage boundaries is based on the following simplifying assumptions: (1) Changes in fluid storage in the confining layer are neglected; (2) confining-layer capacitance effects on heat and solute transport are neglected; (3) flow, heat, and solute transport are affected by the leakage fluxes that enter the region, but flow, heat, and solute conditions that exist on the far side of the confining layer outside the simulation region are not affected by fluxes that enter or leave the simulation region; and (4) flow and transport properties in the confining layer are based on the average of the fluid density and viscosity on either side. These assumptions are quite restrictive; but, in cases where they are not valid, some of the region outside the boundary probably needs to be included in the simulation region. Flow and transport rates are functions of differences in pressure, temperature, and solute-mass fraction at a point in time and are not affected by the previous values of these differences.

## 2.5.3.1. Leaky-Aquifer Boundary

A leaky-aquifer boundary can be adjacent to any part of the simulation region. For illustration, assume that it is part of the upper boundary surface that is overlain by a confining layer. Another aquifer lies above the confining layer with a pressure distribution at its contact with the confining layer which is a known function of time. The geometry is shown in figure 2.2. We are interested in the flux normal to the boundary between the confining layer and the simulation region, located at elevation  $z_B$  in figure 2.2. Under these assumptions, the leakage boundary flux is given by:

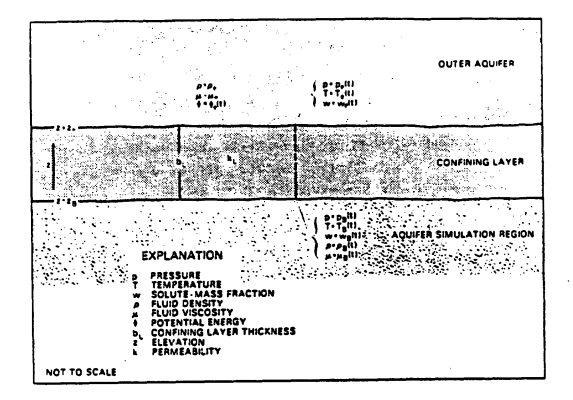


Figure 2.2.--Sketch of geometry for a leaky-aquifer boundary condition.

$$q_{L} = \frac{k_{L}}{\mu_{L}b_{L}} \left[ \rho_{e} \phi_{e} - (p_{B} + \rho_{B}gz_{B}) - (\rho_{e} - \rho_{B})g(z_{e} + z_{B})/2 \right], \text{ for } \underline{x} \text{ on } S^{3}; \qquad (2.5.3.1.1a)$$

with

$$\mathbf{g}_{e} = \frac{\mathbf{r}_{e}}{\rho_{e}} + g z_{e};$$

 $\mu_{L} = \frac{1}{2}(\mu_{B} + \mu_{e});$ 

(2.5.3.1.1b)

(2.5.3.1.1c)

\* where

q<sub>L</sub> is the fluid flux across the leakage boundary (m<sup>3</sup>/m<sup>2</sup>-s).
 Φ<sub>e</sub> is the potential energy per unit mass of fluid in the outer aquifer (N-m/kg);
 p<sub>n</sub> is the pressure at the simulation-region boundary (Pa);

 $p_e$  is the pressure at the top of the confining layer (Pa);  $\rho_B$  is the fluid density at the simulation-region boundary (kg/m<sup>3</sup>);  $\rho_e$  is the fluid density in the outer aquifer (kg/m<sup>3</sup>);  $k_L$  is the permeability of the confining layer (m<sup>2</sup>);  $\mu_L$  is the fluid viscosity in the confining layer (kg/m-s);  $b_L$  is the thickness of the confining layer (m);  $z_B$  is the elevation of the simulation-region boundary (m);  $z_e$  is the elevation at the top of the confining layer (m); and  $S^3$  is the region boundary surface over which a leakage-

boundary condition exists.

The terms  $\Phi_e$ ,  $\rho_e$ ,  $k_L$ ,  $\mu_L$ ,  $b_L$ , and  $z_B$  are specified functions of position along the leakage boundary;  $\Phi_e$  and  $\rho_e$  also can be functions of time. The mass flux is calculated using  $\rho_e$  if the flux is into the simulation region, and using  $\rho_B$  if the flux is out from the simulation region. This choice of density is an approximation because it will take some time for the fluid in the confining layer to attain the limiting value after a change in flow direction takes place. However, this approximation is consistent with the neglect of transient flow and storage effects within the confining layer.

The heat and solute fluxes are assumed to be purely advective. They are obtained from enthalpies and mass fractions of the outer aquifer or at the boundary of the simulation region depending on the flux direction. Thus:

$$q_{HL} = H_e \rho_e q_L$$
, if  $q_L > 0$ , for x on S<sup>3</sup>; (2.5.3.1.2a)  
=  $H_B \rho_B q_L$ , if  $q_L < 0$ ; (2.5.3.1.2b)

$$q_{SL} = w_{\rho} q_{T}, \text{ if } q_{L} > 0, \text{ for } x \text{ on } S^{3};$$
 (2.5.3.1.3a)

$$= w_{\rm p} \rho_{\rm p} q_{\rm f}, \text{ if } q_{\rm f} < 0 ; \qquad (2.5.3.1.3b)$$

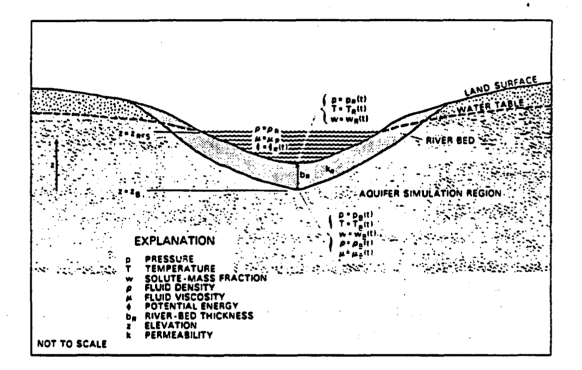
q<sub>HL</sub> is the heat flux across the leakage boundary (W/m<sup>2</sup>);
H<sub>e</sub> is the specific enthalpy of the fluid in the outer aquifer (J/kg);
H<sub>B</sub> is the specific enthalpy of the fluid at the region boundary
 (J/kg);
q<sub>SL</sub> is the solute flux across the leakage boundary (kg/m<sup>2</sup>-s);
w<sub>e</sub> is the solute mass fraction in the outer aquifer (-); and
w<sub>p</sub> is the solute mass fraction at the region boundary (-).

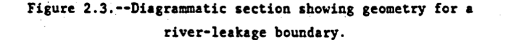
Note that  $H_e$  and  $w_e$  are specified functions of position along the leakage boundary and time.

## 2.5.3.2. River-Leakage Boundary

The river-leakage boundary condition is a second type of leakage-boundary condition that is very similar to a leaky-aquifer condition, with the following differences: (1) This boundary condition is appropriate only for unconfined aquifer regions and is at an upper- or lateral-boundary surface; (2) the less-permeable boundary layer is now the riverbed-sediment layer, that is basically a piecewise-linear feature that traverses the upper boundary of the aquifer region; (3) a limit on the maximum flux from the river to the aquifer is imposed. Additional assumptions for the river-leakage option are: (1) The riverbed thickness is assumed constant over each cross section of the river; and (2) pressure and elevation differences and fluid properties are taken at the river centerline, representing conditions across the riverbed. An area factor is introduced to account for the fact that the riverbed area is only a fraction of the region boundary traversed by the river. The flux limit is set by not allowing the flux to increase after the aquifer pressure plus gravitational potential decreases to less than the gravitational potential at

the bottom of the river. Physically, this means that if the water table declines below the bottom of the riverbed, the increased resistance to flow, because of the porous medium becoming partially saturated, prevents further increases in flux from the river to the aquifer. Thus, the flux limitation is a crude approximation to the physical situation. The simplified geometry of a





river-leakage boundary is shown in figure 2.3. Note that the z-axis is positive in the vertically upward direction. The present version of the HST3D program cannot simulate river leakage with a tilted coordinate system.

With the above assumptions, equations 2.5.3.1.1a and 2.5.3.1.1b become:

$$q_{R} = \gamma_{R} q_{L}; \text{ for } \underline{x} \text{ on } S^{4};$$
 (2.5.3.2.1a)

with

$$\Phi_{e} = g z_{RFS}$$
; (2.5.3.2.1b)

and

$$q_{Rmax} = q_R |_{p_R} = 0;$$
 (2.5.3.2.1c)

where

- $q_R$  is the fluid flux across the river-leakage boundary from the river to the aquifer  $(m^3/m^2-s)$ ;
- $q_{Rmax}$  is the maximum fluid flux from the river to the aquifer  $(m^3/m^2-s)$ ; and

  - $\gamma_R$  is the fraction of riverbed area per unit area of aquifer boundary (-);
  - $z_{\rm RFS}$  is the elevation of the water surface of the river (m); and

S<sup>4</sup> is the region boundary surface over which a river-leakage boundary condition exists.

Note that  $\Phi_e$ ,  $\rho_e$ ,  $k_L$ ,  $b_L$ ,  $\mu_e$ ,  $z_B$ , and  $\gamma_R$  are specified as functions of position along the river length, and at  $\Phi_e$  and  $\rho_e$  also can be functions of time. For calculating  $\Phi_e$ , the value of atmospheric pressure can be taken as zero, because pressures are relative to atmospheric pressure. The mass flux is calculated using  $\rho_e$  if the flux is into the aquifer, and using  $\rho_B$  if the flux is out from the aquifer. The heat and solute fluxes are assumed to be purely advective, and are obtained from the enthalpies and mass fractions of

the river fluid, or from the aquifer at the leakage boundary, depending on the flux direction, as given by equations 2.5.3.1.2a-b and 2.5.3.2.3a-b with  $q_{L}$  replaced by  $q_{R}$ . The enthalpy variation with pressure is neglected or the river-leakage boundary condition, as this variation is assumed to be small.

### 2.5.4. Aquifer-Influence-Function Boundary Conditions

The aquifer-influence-function (AIF) boundary conditions have been presented in the petroleum reservoir-simulation literature. Several methods have been used to calculate water influx at reservoir-aquifer boundaries. For a summary, the reader is referred to Craft and Hawkins (1959, ch. 5) and Aziz and Settari (1979, sec. 9.6).

The utility for ground-water flow simulation of fluid-flux calculations using aquifer-influence functions results from the fact that they enable a simulation region to be embedded within a finite or infinite surrounding region, for which the aquifer properties are known only in a general sense, and where the outer-aquifer-region flow field influences the inner-aquifer region of interest only in a general way. The primary benefit of using AIF boundary conditions is the reduction in size of the simulation region, resulting in a savings in computer-storage requirement and computation time.

Suppose that an aquifer region can be divided into subregions (fig. 2.4), where the inner-aquifer region is the one of primary interest, and the outer-aquifer region is less completely identified with respect to aquifer properties and geometrical configuration. The outer-aquifer region may completely or partially surround the inner-aquifer region, as shown in figures 2.4A and 2.4B. Variable density and nonisothermal flow may be simulated in the inner-aquifer region, but not in the outer-aquifer region. The actual simulation region may be reduced to only the inner-aquifer region, and the boundary condition at the boundary between the two regions (the AIF boundary) is taken to be the AIF boundary condition representing the outer-aquifer region. Aquifer-influence functions are analytical expressions that describe the flow rate, pressure, and cumulative flow at the boundary between the

inner-aquifer and outer-aquifer regions, in response to pressure variations at the boundary. For the purposes of ground-water flow simulation described herein, the cumulative-flow aquifer-influence functions are not of concern. Flow from the outer-aquifer region is assumed to influence the inner-aquifer region of simulation, but flow to the outer-aquifer region does not affect any conditions there.

The aquifer-influence functions that describe transient flow across the AIF boundary are based upon analytical solutions to the ground-water flow equation in the outer-aquifer region. To obtain an analytical solution, the aquifer and fluid properties of the outer-aquifer region are assumed to be constant and uniform, and the geometry of the boundaries between the innerand outer-aquifer regions need to be approximated by simple shapes.

Two types of aquifer-influence functions currently are available for the heat- and solute-transport simulator; one type treats the outer-aquifer region as a "pot;" the other type uses a transient-flow solution for simple-geometry and simple-boundary conditions. An aquifer-influence function based on the assumption of steady-state flow also exists in the petroleum-reservoir simulation literature, but it only is a restricted form of the leakage-boundary condition presented in section 2.5.3.1. Only one type of aquifer-influence function is allowed in any given simulation.

## 2.5.4.1. Pot-Aquifer-Influence Function

The pot-aquifer-influence function is based on the assumption of an outer-aquifer region with exterior boundaries that are impermeable (fig. 2.5). The outer-aquifer region needs to have volume and compressibility that are sufficiently small so that the pressure in this outer-aquifer region always will be virtually in equilibrium with the pressure distribution along the boundary surface between the inner- and outer-aquifer regions. Then, flow will occur in response to the rate of change of pressure at this boundary. The governing equation is obtained from mass conservation in a vertically deforming, compressible, porous medium. Using the Gauss divergence theorem (Karamcheti, 1967, p. 73) and assuming a uniform, constant, fluid density and

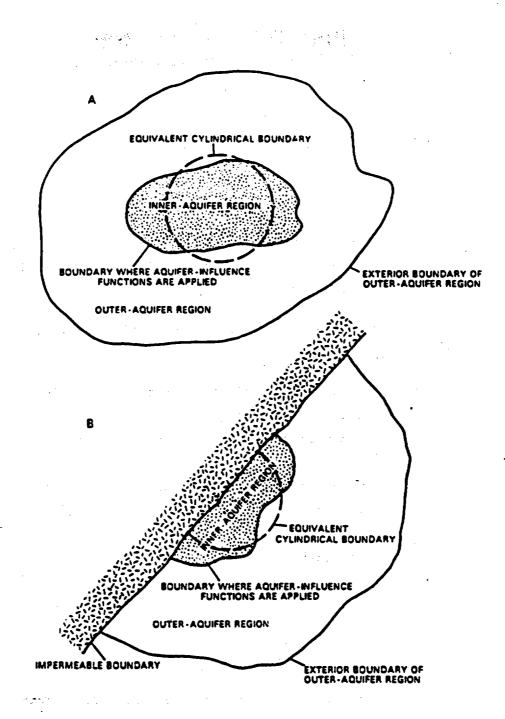


Figure 2.4.--Plan view of inner- and outer-aquifer regions and boundaries: A, Outer-aquifer region completely surrounding inner region; B, Outer-aquifer region half-surrounding inneraquifer region. porosity, we obtain by integration over the outer-region volume:

$$Q_A = (\alpha_{be} + \varepsilon_e \beta_{pe}) \frac{\overline{\partial r_e}}{\partial t} V_e;$$
 (2.5.4.1.1)

#### where

- Q<sub>A</sub> is the volumetric flow rate across the boundary between the inner- and outer-aquifer regions; (positive is into the inner-aquifer region); (m<sup>3</sup>/s).
- $\frac{\partial p_e}{\partial t}$  is the spatial average of the rate of pressure change in the outer region (Pa/s);
- α<sub>be</sub> is the bulk compressibility of the porous medium in the outeraquifer region (Pa<sup>-1</sup>);
- $\underset{\text{pe}}{\overset{\text{is the fluid compressibility in the outer-aquifer region}}{(\text{Pa}^{-1})};$
- $\boldsymbol{\epsilon}_{e}$  is the porosity in the outer-aquifer region (-); and
- V is the volume of the outer-aquifer region  $(m^3)$ .

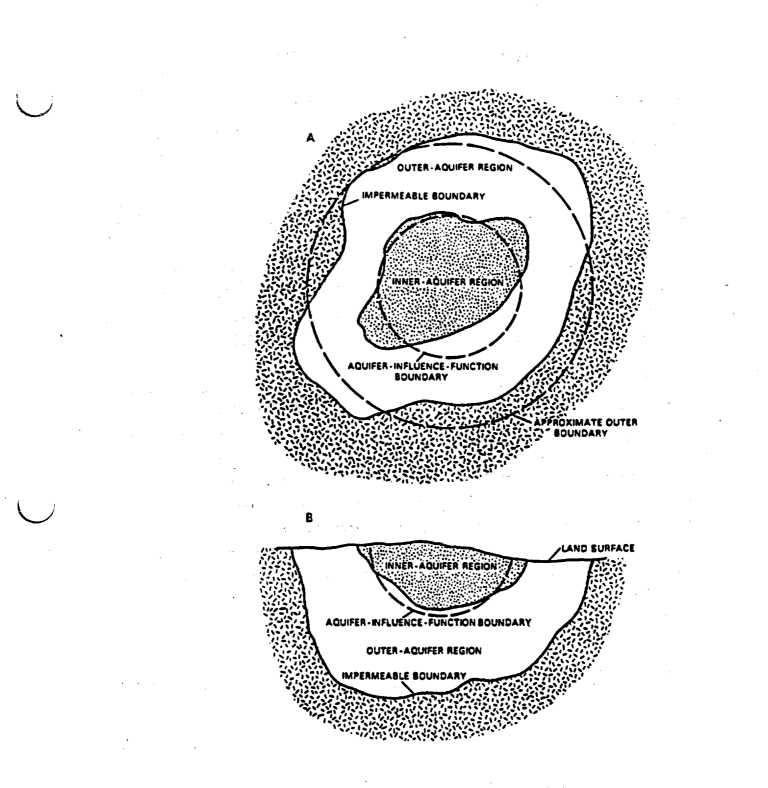
Because pressure equilibrium is assumed:

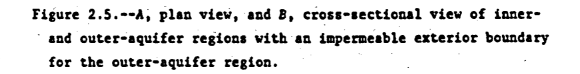
$$\frac{\partial p_e}{\partial t} = \frac{\partial p_B}{\partial t}; \qquad (2.5.4.1.2)$$

where

$$\frac{\partial p_{B}}{\partial t}$$
 is the spatial average rate of pressure change at the boundary of the inner-aquifer region (Pa/s).

Equation 2.5.4.1.1 is in a form suitable for calculating overall fluid flow balances but it is difficult to distribute the flow over the AIF boundary





of the simulation region, particularly when the rate of change of boundary pressure is not uniform over the boundary. Allocation of the flow rate will be explained in section 3.4.4 on numerical implementation.

## 2.5.4.2. Transient-Flow, Aquifer-Influence Function

The transient-flow solution method employs the Carter-Tracy AIF calculation technique (Carter and Tracy, 1960) and analytical solutions presented by Van Everdingen and Hurst (1949). A brief summary of the method follows. A detailed presentation is given by Kipp (1986).

Let the AIF boundary between the inner-aquifer or simulation region and the outer-aquifer region be approximated by a cylinder of a given radius and height. A plan view is presented in figure 2.5A. For a simulation region that is a rectangular prism, this boundary cylinder will be a severe approximation of the actual boundary shape. The outer boundary of this outer-aquifer region is a cylinder at a finite or infinite radius. The thickness of the outer-aquifer region is assumed to be uniform, with impermeable upper and lower boundary surfaces. Ground-water flow in this outer-aquifer region is radial at a given elevation, and the pressure satisfies:

$$(\alpha_{be} + \varepsilon_{e}\beta_{pe}) \frac{\partial p_{e}}{\partial t} = \frac{k_{e}}{\mu_{e}} \left[ \frac{\partial^{2}p_{e}}{\partial r^{2}} + \frac{1}{r} - \frac{\partial p_{e}}{\partial r} \right]; \qquad (2.5.4.2.1)$$

where

r is the radial coordinate (m);
p<sub>e</sub> is the pressure in the outer-aquifer region (Pa);
k<sub>e</sub> is the permeability in the outer-aquifer region (m<sup>2</sup>);
μ<sub>e</sub> is the viscosity in the outer-aquifer region (kg/m-s);
α<sub>be</sub> is the bulk compressibility of the porous medium in the outer-aquifer region (Pa<sup>-1</sup>);

 $\beta_{pe}$  is the compressibility of the fluid in the outer-aquifer region (Pa<sup>-1</sup>); and

 $\varepsilon_{p}$  is the effective porosity of the outer-aquifer region (-).

The initial condition is:

$$at t = 0, p_{a} = p_{a};$$

where

p is the initial uniform pressure (Pa).

Van Everdingen and Hurst (1949) used two different boundary conditions at the AIF cylindrical boundary: one condition was constant pressure, and the other condition was constant flow rate.

The boundary conditions are either specified pressure:

at 
$$r = r_{T}$$
,  $p_{p} = p_{R}$ ;

(2.5.4.2.3a)

where

 $r_{\tau}$  is the interior radius (m); and

 $p_R$  is the constant, specified pressure at the boundary (Pa);

or specified flow rate:

at 
$$r = r_{I}$$
;  $\frac{\partial p_{e}}{\partial r} = \frac{Q_{A} \mu_{e}}{2\pi r_{I} b_{e} k_{e}}$ ;

(2.5.4.2.3b)

(2.5.4.2.2)

where

 $Q_A$  is the constant, specified flow rate at the boundary (positive is from the outer-aquifer region to the inner-aquifer region) (m<sup>3</sup>/s); and

At the exterior cylindrical boundary, the condition is, for an infinite region:

as 
$$r + \infty$$
,  $p_e + p_e$ ; (2.5.4.2.4a)

and, for a finite region, either no flow,

at 
$$r = r_E$$
,  $\frac{\partial p_e}{\partial r} = 0$ ; (2.5.4.2.4b)

where

or specified pressure:

at 
$$r = r_E$$
,  $p_e = p_e^0$ . (2.5.4.2.4c)

Solutions to the dimensionless form of equations 2.5.4.2.1-2.5.4.2.4 are given in Van Everdingen and Hurst (1949) and were derived using Laplace transform techniques. For example, the flow-rate response to a unit change in pressure boundary condition (eq. 2.5.4.2.3a) and the pressure response to a unit withdrawal flow-rate boundary condition (eq. 2.5.4.2.3b) for an infinite outer-aquifer region are:

$$Q_{U}(t') = \frac{4}{\pi^{2}} \int_{0}^{\infty} \frac{e^{-\lambda^{2}t'} d\lambda}{\lambda [J_{0}^{2}(\lambda) + Y_{0}^{2}(\lambda)]};$$
 (2.5.4.2.5a)

and

$$P_{U}(t') = \frac{-4}{\pi^{2}} \int_{0}^{\infty} \frac{1 - e^{-\lambda^{2} t} d\lambda}{\lambda^{3} [J_{1}^{2}(\lambda) + Y_{1}^{2}(\lambda)]}; \qquad (2.5.4.2.5b)$$

respectively, with

$$t' = \frac{k_e t}{r_I^2 (\alpha_{be} + \epsilon_e \beta_{pe}) \mu_e};$$
 (2.5.4.2.5c)

where

 $J_i$  is the Bessel function of the first kind of order i;  $Y_i$  is the Bessel function of the second kind of order i; and t' is the dimensionless time (-).

Equation 2.5.4.2.5a was presented by Jacob and Lohman (1952) in a different form as a solution to the constant drawdown problem for flow to a well. These two aquifer-influence functions will be referred to as the flow-rate response to a unit-step pressure change  $Q_U^{\prime}$ , and the pressure response to a unit-step withdrawal flow rate,  $P_{\rm H}^{\prime}$ , respectively.

The concept of superposition or convolution (Tychonov and Samarski, 1964, p. 209) is used to derive the aquifer-influence functions from the unit-step response functions for a transient-pressure function at the boundary between the inner-and outer-aquifer regions. Thus, the flow-rate response of this ground-water flow system to a time-varying pressure at the inner boundary of the outer-aquifer region can be written as:

and at

$$z_n = b_{HC}; T_e = T_e^{0}(b_{HC});$$
 (2.5.5.1d)

where

 $T_{p}$  is the boundary temperature at the aquifer boundary (°C),

b<sub>HC</sub> is the effective thickness of the conducting medium outside the region (m).

The thermal properties of the adjacent medium are assumed constant and uniform. Thus:

$$D_{He}^{} = \frac{k_{e}}{\rho_{se}c_{se}};$$
 (2.5.5.2)

where

 $D_{\mu_0}$  is the thermal diffusivity for the adjacent medium (m<sup>2</sup>/s).

Since the heat flux depends on the temperature profile in the exterior medium which in turn depends on the thermal history of the simulation, a simplifying approximation is used. This approximation eliminates the need to recompute or save the temperature-profile history during the course of the simulation.

The boundary-value problem specified by equations 2.5.5.1a-d can be resolved into simpler problems, as shown by Sneddon (1951, p. 162-165) or Tychonov and Samarski (1964, p. 203-209), using various forms of Duhamel's Theorem. Two simpler problems, for a general time interval, are:

$$\frac{\partial T_1}{\partial t} = D_{\text{He}} \frac{\partial^2 T_1}{\partial z_n^2}; \quad \text{on } 0 \le z_h \le b_{\text{HC}} \text{ and } t_o \le t \le t_1 . \quad (2.5.5.3a)$$

Boundary conditions:

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at 
$$z_n = 0$$
;  $T_1 = 0$ ; (2.5.3b)

$$t z_n = b_{HC}; T_1 = 0.$$
 (2.5.3c)

Initial condition:

at 
$$t = t_0$$
;  $T_1 = T_e^0(z_n)$ ; (2.5.5.3d)

and

$$\frac{\partial T_2}{\partial t} = D_{\text{He}} \frac{\partial^2 T_2}{\partial z_n^2} \quad \text{on } 0 \le z_n \le b_{\text{HC}} \text{ and } t_0 \le t \le t_1 \quad (2.5.5.4a)$$

Boundary conditions:

at 
$$z_n = 0$$
;  $T_2 = T_B(t)$ ; (2.5.5.4b)

at 
$$z_n = b_{HC}$$
;  $T_2 = 0$ . (2.5.5.4c)

<sup>\*</sup>Initial condition:

at 
$$t = t_0$$
;  $T_2 = 0$ ; (2.5.5.4d)

where

-)

 $T_1(z,t)$  is the temperature solution to the first heat-conduction problem (°C); and

 $T_2(t)$  is the temperature solution to the second heat-conduction problem (°C).

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The total temperature solution is the sum of  $T_1$  and  $T_2$  and the boundary heat flux is derived from the gradient of this temperature. However, because the boundary-temperature function is not known in advance the following approximation is made

$$q_{HC} = -K_{e} \frac{\partial T_{e}}{\partial z_{n}} \begin{vmatrix} 2.5.5.5a \\ Z_{n} = 0 \\ T_{2} = T_{B}(t_{o}) \end{vmatrix}$$
(2.5.5.5a)

$$= -K_{e} \begin{bmatrix} \frac{\partial(T_{1}+T_{2})}{\partial z_{n}} \\ z_{n} \end{bmatrix} z_{n} = 0 \\ T_{2}=T_{B}(t_{0})$$

$$\begin{array}{c|c} + \frac{\partial}{\partial T_{B}} & \frac{\partial(T_{1}+T_{2})}{\partial z_{n}} & \delta T_{B} \\ & z_{n}=0 \\ & T_{2}=T_{B}(t_{0}) \end{array}; \qquad (2.5.5.5b)$$

where

 $q_{HC}$  is the heat flux at a heat-conduction boundary at a given boundary temperature and time (W/m<sup>2</sup>); and  $\delta T_B$  is the change in boundary temperature in the time interval t<sub>o</sub> to t (°C).

Equation 2.5.5.5b is simply a Taylor-series expansion of the flux as a function of the variable boundary temperature.

By interchanging the order of differentiation and using the facts that

$$\frac{\partial T_1}{\partial T_p} = 0 \quad ; \tag{2.5.5.6a}$$

(2.5.5.6b)

and

$$\frac{\partial T_2}{\partial T_B} = T_U(t);$$

where

 $T_{U}$  is the solution to equations 2.5.5.4a-d with  $T_{B} = 1$  (°C);

we obtain:

$$q_{HC} = -K_{e} \begin{bmatrix} \frac{\partial T_{e}}{\partial z_{n}} & \frac{\partial T_{u}}{\partial z_{n}} & \frac{\delta T_{B}}{\partial z_{n}} \\ T = T_{B}(t_{0}) \end{bmatrix}, \text{ for } \underline{x} \text{ on } S^{5}; \quad (2.5.5.7)$$

where

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 $S^5$  is the part of the boundary that is a heat-conduction boundary.

The temperature, T, now satisfies equations 2.5.5.1a-d with the time dependence of the boundary condition removed in equation 2.5.5.1c. This approach to the treatment of heat-conduction boundary conditions was presented by Coats and others (1974) in the appendix to their paper. A heat-conduction boundary condition also could be treated like the transient AIF boundary condition, but that is beyond the scope of this work.

## 2.5.6. Unconfined Aquifer, Free-Surface Boundary Condition

For an unconfined aquifer, a free-surface boundary exists with a position in space and time that is unknown before the flow equations are solved. Therefore, two boundary conditions need to be imposed. The first is that pressure is atmospheric at the free surface. The second is the kinematic condition expressing the fact that the movement of this surface of atmospheric pressure needs to satisfy a continuity equation at the free surface.

The free-surface boundary is assumed to be a sharp interface between the fully saturated region of simulation and the unsaturated porous medium outside. The zone of capillary fringe that is partially saturated and the surfaces of seepage that exist with free-surface gravity flow are neglected. Delayed yield effects also are neglected, therefore, the specific yield is equal to the effective porosity,  $\varepsilon$ , in the vicinity of the free surface. The effective porosity under draining conditions is less than the porosity used to calculate interstitial velocity (Bear, 1972, p. 255) but the difference is assumed negligible for the HST3D simulator. Finally, the z-axis is assumed to point vertically upward when an unconfined aquifer is being simulated.

The heat- and solute-transport simulator treats the free-surface boundary in an approximate fashion. The approach follows the ideas of Prickett and Lonnquist (1971, p. 43-45) extended to a three-dimensional flow and variable-density system. The pressure condition:

$$p = 0$$
, on  $S^{6}(x,t)$ ; (2.5.6.1)

where

S<sup>6</sup> is the free-surface location that varies in space and time; is employed, but the kinematic boundary condition is neglected. The absolute pressure on the free surface is atmospheric, so the relative pressure is zero. Hydrostatic conditions are assumed to exist in the immediate vicinity of the free surface. The location of the free surface is determined by interpolation in the calculated pressure field to determine the location where equation 2.5.6.1 is satisfied. Under this approximate treatment, the free surface

moves in response to a net gain or loss of fluid in its vicinity. Thus, fluid mass is conserved, but the kinematics of the free-surface movement are neglected. This approximation is acceptable when the velocity of free-surface movement is small relative to the horizontal interstitial velocity.

In addition, the computational region is fixed for the duration of the simulation. Boundary pressures less than atmospheric imply that the free surface is below the boundary of the region; whereas boundary pressures greater than atmospheric imply that the free surface is above this boundary. As will be explained in section 3.4.6, the free surface is allowed to rise above the region boundary a short distance which is a function of the vertical discretization. This allowance enables the free surface to move within a reasonable range during a simulation. The fluid- and porous-matrix compressibilities usually are taken to be zero for unconfined flow systems, and the user may specify compressibility values of zero for the HST3D simulator.

# 2.6. INITIAL CONDITIONS

This heat- and solute-transport simulation code solves only the transient forms of the ground-water flow and the two transport equations, thus initial conditions are necessary to begin a simulation. Several options are available.

For the flow equation, an initial-pressure distribution within the region needs to be specified. This can be done as a function of position or can be set to hydrostatic conditions, with the pressure given at one elevation. In the case of nearly uniform and constant density, an initial potentiometrichead distribution can be specified, which is the water-table elevation. The water-table elevation is specified for the upper layer of cells only. No option to specify a velocity field as an initial condition exists.

For the heat-transport equation, the initial-temperature field needs to be specified. Again, this can be done as a function of position, or interpolated along the z-coordinate direction from a specified geothermal profile.

For the solute-transport equation, the initial mass-fraction field needs to be specified. This can be done only by specifying values as a function of position.

As will be described in section 4.6, pressure, temperature, and massfraction fields calculated by one simulation can be used as the initial conditions for another simulation, using the restart option. This often is the easiest way to establish a steady-state flow field before transport is simulated. Of course, one needs to determine whether or not an initial steady-state flow field exists for the physical situation being simulated. It should be noted that, with a hydrostatic or other estimate of initial pressure conditions, it could take some time to establish the steady-state flow field.

Mathematically, the initial conditions can be stated as follows: At t=0:

$p = p^{o}(x)$ , in V;	(2.6.1a)
$T = T^{0}(x)$ , in V;	(2.6.1b)

 $w = w^{0}(x), \text{ in } V;$  (2.6.1c)

where

<u>x</u> is the vector of position (m), V is the simulation region; and p<sup>0</sup>, T<sup>0</sup>, w<sup>0</sup> are the initial dependent variable distributions (Pa,°C,-). In order to perform numerical calculations that solve the governing equations, we first need to discretize the partial-differential equations and boundary condition relations in space and time. Various algorithms are used to determine parameters and to implement the boundary conditions. Then the flow and two transport equations are solved sequentially after they have been modified by a partial Gauss reduction. Finally, the sets of discretized equations are solved repeatedly, as the simulation time advances, using a direct or an iterative equation solver. This chapter will cover each of these steps for the numerical-simulation calculation.

## 3.1 EQUATION DISCRETIZATION

The classical method of finite differences is used to discretize the partial-differential equations and boundary conditions in space and time. Several options are available for the differencing.

The first step in spatial discretization is to construct a mesh or grid of node points and their associated cells, that covers the simulation region to a close approximation (fig. 3.1). The grid of node points is formed by specifying the distribution of nodes in each of the three coordinate directions; (two directions, if a cylindrical-coordinate system). The volume associated with each node will be called a cell; it is formed by the cell boundaries, which are planes that bisect the distance between adjacent node points. Thus, for the case of unequal nodal spacing, the node points do not lie at the centers of their respective cells. Boundaries are represented by planes containing node points. Thus, half-, quarter-, and eighth-cells appear at various sides, edges, and corners of the mesh, forming the simulation region (fig. 3.1). The minimum number of nodes required to define a region is eight, one node at each corner of the rectangular prism. The mesh or grid described is called a point-distributed grid. Other terms that have been used are face-centered mesh and lattice-centered mesh or grid. Another term that has been used for cell is block.

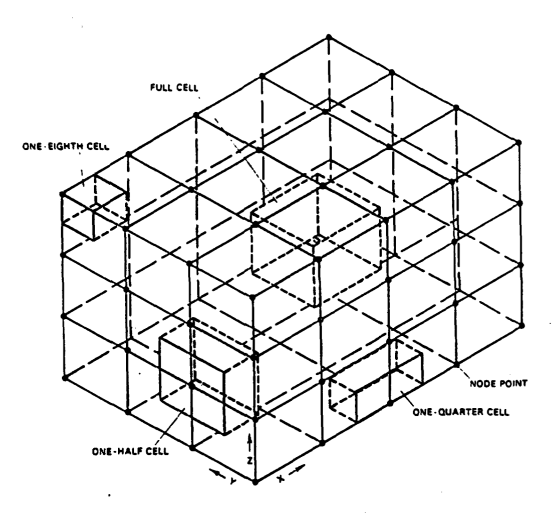
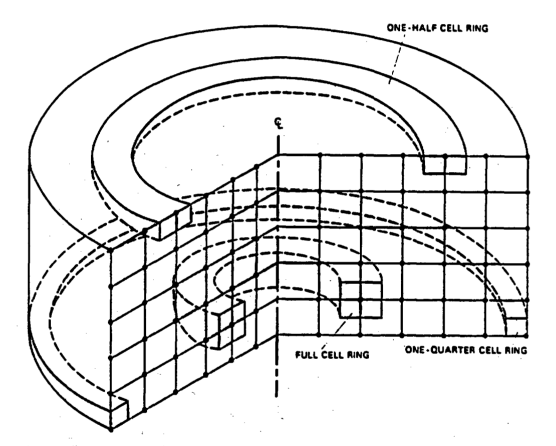
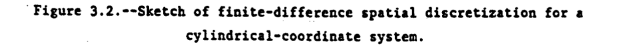


Figure 3.1.--Sketch of finite-difference spatial discretization of the simulation region.

The simulation region is discretized into rectangular prisms for the cartesian-coordinate case (fig. 3.1) and into annuli with rectangular cross sections for the cylindrical-coordinate case (fig. 3.2). Four types of regional volume subdivisions are defined (fig. 3.3). The primary subdivision is the cell that is the volume over which the flow, heat, and solute balances

are made to give the nodal finite-difference equations. The second subdivision is the element that is the volume bounded by eight corner nodes in cartesian coordinates and four corner nodes in cylindrical coordinates. The element is the minimum volume with uniform porous-medium properties. The third subdivision is the zone that is a continuous set of elements with the





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same porous-medium properties. The one restriction is that zones need to be convex. In other words, they need to be rectangular prisms. One zone may not border another zone on more than one side. Sometimes multiple, adjacent zones will have to be specified that have the same properties in order to adhere to this restriction of convex shape. The fourth subdivision is the subdomain that is the intersection or common volume of an element with a cell. A cell may have as many as eight subdomains, if it is an interior cell, or as few as one subdomain, if it is a corner cell. The finite-difference equations are

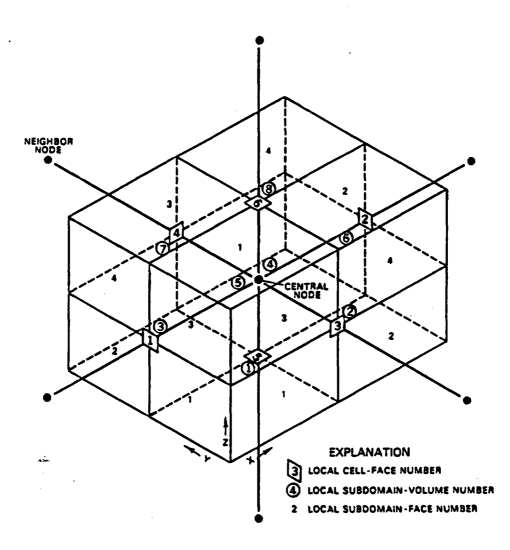


Figure 3.3.--Sketch of a node with its cell volume showing the cell faces, the subdomain volumes, and the subdomain faces.

assembled by adding the contributions of each subdomain in turn to the equation for a given cell. The primary reason for introducing the concepts of elements and zones for assigning porous-medium properties is so that porous-medium properties can be defined easily at the cell boundaries without the need for harmonic-mean calculations.

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A numbering scheme local to the cell is used during coefficient calculation and assembly. Figure 3.3 shows the subdomain local numbering from one to eight. The six faces are numbered as shown in figure 3.3 and each face is subdivided into four subdomain faces with numbers for the visible faces as shown. For the cylindrical system, the corresponding volumes and faces are numbered in figure 3.4.

A common alternative method for constructing the mesh is to specify the locations of the planes that form the cell walls. Their intersections form the cells; then the node points are located in the center of each cell. This is called a cell-centered or block-centered grid. One advantage of this grid is that fewer cells are required to span a given simulation region, because fractional cells do not appear at the boundaries.

The point-distributed grid was selected for this simulator, because the finite-difference spatial approximations to the dispersive terms in the flow and transport equations are consistent and convergent for the pointdistributed grid under conditions of variable-grid spacing; whereas, these approximations are not necessarily consistent and convergent for the cellcentered grid. As shown by Settari and Aziz (1972, 1974), the local truncation error for the cell-centered grid has a term that does not necessarily vanish as the grid spacing is refined. A second reason for selecting the point-distributed grid is that the presence of nodes on the boundary surfaces simplifies the treatment of certain boundary conditions. It is common to approximate spatially distributed, aquifer properties as uniform zones. A disadvantage of the point-distributed grid is that it is difficult to locate the cell boundaries, so that they coincide with the zoned-property boundaries. This difficulty can be avoided by making the properties uniform over an element rather than a cell. This will be described in the parameterdiscretization section.

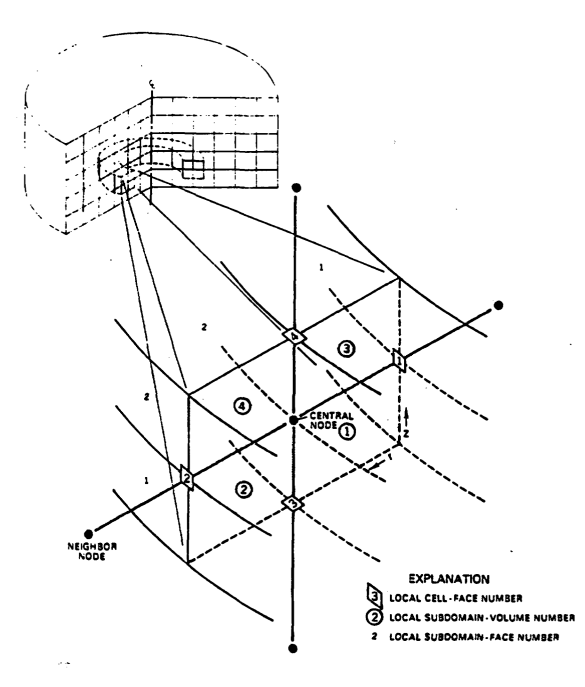


Figure 3.4.--Sketch of a node in a cylindrical-coordinate system with its cell volume, showing the cell faces, the subdomain volumes, and the subdomain faces. A cartesian-coordinate grid for a region is shown in figure 3.1. Note that the basic cell is a rectangular prism; thus, region boundaries that are not parallel to a coordinate axis must be approximated by a staircase-like pattern of cell boundaries. No provision for boundary faces that are diagonally oriented to the coordinate axes exists in the present version of the HST3D code. It can also be seen that the entire simulation region needs to be contained in a large rectangular prism. The nodal dimensions of this prism are the maximum number of nodes along the three coordinate axes,  $N_x$ ,  $N_y$ , and  $N_z$ . Approximation of diagonal boundaries by a staircase-like pattern will cause a set of cells to be within the large prism that are excluded from the simulation region. These cells will be referred to as excluded cells.

The method used for spatial discretization is a subdomain weightedresidual method with approximating functions that are piecewise linear for the dependent variables. The unknown parameters in the approximating functions are the nodal values of the dependent variables. The residuals are the errors in the governing equations that result from using approximating functions to the exact solutions, and equations for the unknown parameters require that the average residual over each cell is zero (Crandall, 1956, p. 149; Finlayson, 1972, p. 7-9, 137, 142). The partial-differential equations are discretized in space by integrating them over each cell volume. Then the divergence theorem of Gauss (Karamcheti, 1967, p. 73) is used to transform the volume integrals of the divergence terms into surface integrals of a normal derivative. The spatial derivatives in the surface integrals are approximated by central or upstream differences. The volume integrals are approximated easily, using the mean value theorem of integral calculus, because all fluid and porous medium properties are assumed to be constant throughout the subdomain volumes of a cell. For the integral of the time derivative, we assume that the time derivative evaluated at the node approximates the spatial average of the derivative over the volume of the cell. Thus, the capacitancecoefficient matrix of the temporal derivative terms is diagonalized. This method for spatial discretization is conceptually similar to the integratedfinite-difference method presented by Narasimhan and Witherspoon (1976).

The porous-matrix hydraulic, thermal-transport, and solute-transport properties are discretized on an element basis, with a set of elements forming a zone of constant properties. The dependent variables that are properties of the fluid are discretized on a cell basis. Boundary-condition fluxes and source flow rates also are discretized on a cell basis.

To illustrate the discretization of the flow, heat, and solute equations, we shall use a general transport equation. The procedure follows that of Varga (1962, sec. 6.3), Spanier (1967, p. 218-222), Cooley (1974, p. 10-13) or Roache (1976, p. 23-28) extended to include spatial first-derivative terms, to three dimensions for cartesian coordinates, and to handling dispersive tensors that are not necessarily diagonal. The restriction exists that all cellboundary planes need to be perpendicular to a coordinate direction. The general transport equation has the form of a parabolic, partial-differential equation:

$$\frac{\partial}{\partial t} (A(\underline{x},t) u(\underline{x},t)) = \nabla \cdot \underline{B}(\underline{x},t) \cdot \nabla u(\underline{x},t) - \nabla \cdot \underline{C}(\underline{x},t) u(\underline{x},t)$$
(3.1.1a)

$$N_{s} + D(\underline{x},t)u(\underline{x},t) + \sum_{s=1}^{N} E_{s}(t)\delta(\underline{x}-\underline{x}_{s});$$

where

x is the vector of position, (x,y,z), (m); A is the capacitance coefficient (appropriate units); B is the tensor of diffusion or dispersion of rank 3 (appropriate units); C is the vector of interstitial velocity (m/s); D is the source factor for chemical reaction (appropriate units); E is the source-term intensity (appropriate units); N is the number of source terms; u is the dependent variable (appropriate units); and ô(x-x\_) is the delta function for a point source at x=x\_ (-). Initial condition is:

$$t = 0; u = u^{0}(\underline{x});$$
 (3.1.1b)

Boundary conditions are:

specified value:

$$u = u_{p}(\underline{x}, t), \text{ for } \underline{x} \text{ on } S^{1};$$
 (3.1.1c)

specified flux:

$$-\underline{\beta}(\underline{x},t) \frac{\partial u}{\partial n} + \underline{C}(\underline{x},t) \cdot \underline{n} u = \underline{J}(\underline{x},t) \cdot \underline{n} \text{ for } \underline{x} \text{ on } S^2 ; \qquad (3.1.1d)$$

where

u<sup>0</sup> is the initial distribution of u;

 $u_{R}$  is the boundary distribution of u;

is the derivative in the direction of the outward normal at the on boundary;

 $\underline{J} \cdot \underline{n}$  is the specified total flux normal to the boundary surface; and  $\underline{C} \cdot \underline{n}$  is the advective flux normal to the boundary surface.

#### 3.1.1 Cartesian Coordinates

Integration of equation 3.1.1a over the cell volume associated with a mesh point, m, at a given location with indices i, j, k, gives:

$$\int_{\nabla_{m}}^{\infty} \frac{\partial}{\partial t} Au \, dV = \int_{\nabla_{m}}^{\infty} \nabla \cdot (\underline{\beta} \cdot \nabla u) \, dV - \int_{\nabla_{m}}^{\infty} \nabla \cdot \underline{C} u \, dV \qquad (3.1.1.1)$$

+ 
$$\int_{V_m} \frac{D}{D} u dV + \Sigma \int_{V_m} \frac{E\delta(\underline{x}-\underline{x}_s) dV}{s=1} dV;$$

where

$$V_{m}$$
 is the volume of cell m.

Now, using the previously stated assumption about the integral of the time derivative:

$$\int_{V_{m}} \frac{\partial}{\partial t} \operatorname{Aud} V = \frac{\partial}{\partial t} \int_{V_{m}} \operatorname{Aud} V. \qquad (3.1.1.2)$$

Use of the Gauss divergence theorem on the dispersive and advective terms yields:

$$\int_{V_m} \nabla \cdot (\underline{\underline{a}} \cdot \nabla \underline{u}) dV = \int_{S_m} (\underline{\underline{a}} \nabla \underline{u}) \cdot \underline{\underline{n}} dS ; \qquad (3.1.1.3)$$

and

$$\int_{V_{\underline{m}}} \nabla \cdot (\underline{C}\underline{u}) dV = \int_{S_{\underline{m}}} (\underline{C}\underline{u}) \cdot \underline{n} dS ; \qquad (3.1.1.4)$$

where

S is the boundary of cell m; and <u>n</u> is the outward unit normal vector to the boundary.

Then equation 3.1.1.1 becomes:

$$\frac{\partial}{\partial t} \int_{V_{m}} AudV = \int_{S_{m}} [\underline{\underline{B}} \cdot \nabla u - \underline{\underline{C}} u] \cdot \underline{\underline{n}} dS + \int_{V_{m}} \underline{\underline{D}} udV +$$

$$\int_{V_m} E_{s(m)} \delta(\underline{x} - \underline{x}_{s(m)}) dV . \qquad (3.1.1.5)$$

We have consolidated all the line sources within cell m into a single equivalent line source, thus eliminating the summation. Since considerable arbitrariness exists in selecting the finite-difference approximation of equation 3.1.1.5, we shall choose finite differences that preserve the conservation of u for each cell.

Following Varga, 1962, p. 253, or Cooley, 1974, p. 16, we approximate the rate of change of u in the cell using the mean-value theorem giving:

$$\frac{\partial}{\partial t} \int_{V_m} AudV \cong \frac{\partial}{\partial t} (u(\underline{x}_m, t) \int_{V_m} AdV); \qquad (3.1.1.6)$$

where

 $\underline{x}_m$  is the vector of the node point location (m).

This approximation diagonalizes the coefficient matrix of the temporalderivative terms. The value of the dependent variable at the node is taken to represent the average value over the cell. Now each cell may consist of up to eight subdomains, as shown in figure 3.3, and each subdomain may have different spatial properties. Thus, the integral of A in equation 3.1.1.6 is actually:

$$\begin{array}{c} 8 \\ \Sigma \\ s=1 \end{array} \begin{array}{c} 8 \\ ms \end{array} \begin{array}{c} 8 \\ s=1 \end{array} \begin{array}{c} 8 \\ ms \end{array} \begin{array}{c} \\ s=1 \end{array} \begin{array}{c} 8 \\ ms \end{array} \begin{array}{c} \\ s=1 \end{array} \begin{array}{c} (3.1.1.7) \end{array}$$

where

A is the value of A in subdomain s of cell m; and V is the volume of subdomain s of cell m  $(m^3)$ .

The dispersive-flux term is approximated, recognizing that the surface of cell m is composed of six faces, and that each face belongs to four elements, each of which may have different spatial properties (fig. 3.3). Thus:

$$\int_{S_{m}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS = \Sigma \sum \int_{S_{m}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS ; \qquad (3.1.1.8)$$

where

S is the part of the cell surface that belongs to face p in element  $q (m^2)$ .

A typical integral over a cell face is of the form, for p = 2, as an example:

$$\int_{S_{m2}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS = \sum_{q=1}^{4} \int_{S_{m2q}} [B_{xx} \frac{\partial u}{\partial x} + B_{xy} \frac{\partial u}{\partial y} + B_{xz} \frac{\partial u}{\partial z}] dS ; (3.1.1.9)$$

where

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 $B_{ij}(t)$  are the tensor components of  $\underline{B}$  for a face whose outward normal points in the ith direction, i=x,y, or z.

A sample subdomain volume for subdomain s=1 in figure 3.3 is:

$$V_{1} = \frac{1}{2} (x_{i} - x_{i-1}) \frac{1}{2} (y_{j} - y_{j-1}) \frac{1}{2} (z_{k} - z_{k-1})$$
(3.1.1.10)

A sample cell-face area belonging to face p=2 and zone q=1 in figure 3.3

$$S_{m21} = \frac{1}{2} (y_j - y_{j-1}) \frac{1}{2} (z_k - z_{k-1})$$
(3.1.11)

Equation 3.1.1.9 is based on the fact that each cell face has a normal vector that is alined with one of the cartesian-coordinate directions. Note that  $B_{ij}$  is assumed to be spatially constant over the element q. The partial derivatives are approximated by central differences across each face. Thus, for the face midway between  $x_i$  and  $x_{i+1}$ , denoted by p=2, the outward normal is in the positive x-direction. An integral over this cell face becomes:

$$\int_{S_{m2}} (\underline{B} \cdot \nabla) \cdot \underline{n} dS \cong \sum_{q=1}^{4} \left[ B_{xx}(q) \frac{\partial u}{\partial x} \right|_{i+\frac{1}{2},j,k} S_{m2q}^{+} B_{xy}(q) \frac{\partial u}{\partial y} \right|_{q} S_{m2q}^{-}$$

 $+B_{xz}(q)\frac{\partial u}{\partial z} g_{m2q}$ ; (3.1.1.12)

where

is:

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 $\frac{\partial u}{\partial x} \begin{vmatrix} i+\frac{1}{2}, j, k \end{vmatrix}$  is the gradient of u in the x-direction across the p=2 face at  $y_j$ ,  $z_k$ ;

 $\frac{\partial u}{\partial y}$  is the gradient of u in the y-direction for the subface in the qth element; and

 $\frac{\partial u}{\partial z}$  is the gradient of u in the z-direction for the subface in the qth element. Now

$$\frac{\partial u}{\partial x} |_{i+\frac{1}{2},j,k} \cong \frac{\frac{u_{i+1,j,k} - u_{i,j,k}}{x_{i+1} - x_{i}}; \qquad (3.1.1.13)$$

where

$$u_{i,j,k}$$
 is the value of u at node  $x_i, y_j, z_k$ .

The approximation for  $\frac{\partial u}{\partial y}$  depends on q. For example, for the element bounded by the planes at  $x_i$  and  $x_{i+1}$ ,  $y_j$  and  $y_{j+1}$ ; and  $z_k$  and  $z_{k+1}$ ; denoted by p=2, q=4 in figure 3.3, we have:

$$\frac{\partial u}{\partial y}\Big|_{q=4} \cong \frac{\frac{u_{i+\frac{1}{2},j+1,k} - u_{i+\frac{1}{2},j,k}}{y_{j+1} - y_{j}}}{(3.1.1.14)}$$

where

$$u_{i+\frac{1}{2},j,k} = \frac{1}{2}(u_{i+1,j,k} + u_{i,j,k}).$$
 (3.1.1.15)

Similarly,

$$\frac{\partial u}{\partial z} |_{q=4} \cong \frac{\frac{u_{i+\frac{1}{2},j,k+1} - u_{i+\frac{1}{2},j,k}}{z_{k+1} - z_{k}}.$$
 (3.1.1.16)

The advective-transport term is treated in a similar fashion, but it is somewhat simpler. First, we break it into a sum over the faces and zones:

$$\int_{S_{m}} (\underline{C}u) \cdot \underline{n} dS = \Sigma \Sigma \int_{S} (\underline{C}u) \cdot \underline{n} dS . \qquad (3.1.1.17)$$

A typical integral over a cell face, p=2, is:

$$\int_{S_{m2}} (\underline{C}u) \cdot \underline{n} dS = \sum_{q=1}^{4} \int_{S_{m2q}} C_x u dS ; \qquad (3.1.1.18)$$

where

 $C_{\rm X}$  is the vector component of <u>C</u> for a face whose outward normal points in the x-direction.

Now if the integral is approximated, for the same example face as above, p=2, by:

$$\int_{S_{m2}} C_{x} udS \cong \sum_{q=1}^{L} C_{x}(m,2,q)^{L} (u_{i+1,j,k} + u_{i,j,k}) S_{m2q}; \qquad (3.1.1.19)$$

where  $C_x$  (m,2,q) is the value of  $C_x$  on the face p=2 in element q.

This will lead to a central difference for the advective term of equation 3.1.1.5. If, instead, the following approximation is used:

$$\int_{S_{m2}} C_{x} udS \cong \sum_{q=1}^{4} C_{x}(m,2,q) u_{i,j,k} S_{m2q}, \text{ for } C_{x} > 0 ; \qquad (3.1.1.20a)$$

or

. . . . .

$$\stackrel{4}{\cong} \sum_{q=1}^{C} C_{x}(m,2,q) u_{i+1,j,k} S_{m2q}, \text{ for } C_{x} < 0; \qquad (3.1.1.20b)$$

this will lead to an upstream difference for the advective term. Central differencing may produce oscillations in the solution, whereas upstream

differencing cannot (Price and others, 1966; Roache 1976, p. 161-165). But the penalty to eliminate oscillation resulting from spatial differencing is the addition of artificial dispersion (Roache, 1976, p. 64-66; Lantz, 1970), which can be regarded as smearing out of steep concentration or temperature gradients caused by the numerical method rather than the dispersive mixing term.

An approximation has the transportive property, if a disturbance in the field of property u, is advected only in the direction of the velocity. Recall that C represents the velocity in these equations. The central approximation of equation 3.1.1.19 does not have the transportive property, whereas the upstream approximation of equations 3.1.1.20a and 3.1.1.20b does. However, not all upstream approximations have the transportive property (Roache, 1976, p. 69). While the transportive property is desirable on physical grounds, the grid spacing must be limited to avoid excessive artificial dispersion caused by the numerical method. The criteria for avoidance will be presented in a later section. The numerical implementation of the heat- and solute-transport simulator offers the choice of central or upstream differencing for the advective terms. If upstream differencing is selected, the user must determine the grid spacing that limits numerical dispersion to an acceptable amount.

The source term in equation 3.1.1.5 that is linearly proportional to the value of the dependent variable, u, is averaged throughout the cell volume to obtain the finite-difference approximation. The mean-value theorem is used to approximate the integral, with  $\underline{x}_{m}$  being the node-point location. The volume integral is split into the contributions from the eight subdomains. Thus:

$$\int_{V} Du \, dV \cong u(\underline{x}_{m}, t) \sum_{s=1}^{8} \int_{V} D(s) dV; \qquad (3.1.1.21a)$$

$$\begin{array}{c} & & & & \\ \Xi & u_{i,j,k} & & \Sigma D & V_{i,j,k} \\ & & s=1 \\ \end{array}$$
 (3.1.1.21b)

where

}

 $D_{mS}$  is the value of D in subdomain s of cell m.

The source term at the end of equation 3.1.1.5 is assumed to be a line source in the z-direction of constant intensity, that fully penetrates the cell at  $x=x_s$ ;  $y=y_s$ . Discretization is achieved by carrying out the integration. Thus:

$$\int_{V_m} E_{sm} \delta(x-x_s) \delta(y-y_s) dV = E_m . \qquad (3.1.1.22)$$

This shows that a line source becomes distributed throughout the cell volume, and the precise location is lost in the finite-difference equation.

Combining equations 3.1.1.6 through 3.1.1.22 gives the finite-difference approximation to equation 3.1.1.1a for an interior node or cell. It is of the form:

$$\frac{\partial}{\partial t} (a_{28}u_{i,j,k}) = a_1 u_{i-1,j-1,k-1} + a_2 u_{i,j-1,k-1} + a_3 u_{i+1,j-1,k-1} + a_4 u_{i-1,j,k-1} + a_5 u_{i,j,k-1} + a_6 u_{i+1,j,k-1} + a_7 u_{i-1,j+1,k-1} + a_8 u_{i,j+1,k-1} + a_9 u_{i+1,j+1,k-1} + a_{10} u_{i-1,j-1,k} + a_{11} u_{i,j-1,k} + a_{12} u_{i+1,j-1,k} + a_{13} u_{i-1,j,k} + a_{14} u_{i,j,k} + a_{15} u_{i+1,j,k} + a_{16} u_{i-1,j+1,k} + a_{17} u_{i,j+1,k} + a_{18} u_{i+1,j+1,k} + a_{19} u_{i-1,j-1,k+1} + a_{20} u_{i,j-1,k+1} + a_{21} u_{i+1,j-1,k+1} + a_{22} u_{i-1,j,k+1} + a_{23} u_{i,j,k+1} + a_{24} u_{i+1,j,k+1}$$

+  $a_{25}$   $u_{i-1,j+1,k+1}$  +  $a_{26}$   $u_{i,j+1,k+1}$  +  $a_{27}$   $u_{i+1,j+1,k+1}$ 

It is important to observe that the dependent variable for each interior node is related to 26 other nodal values of that variable, through the finitedifference equation in space. The six nearest neighbors to a given node appear in the terms with coefficients  $a_5$ ,  $a_{11}$ ,  $a_{13}$ ,  $a_{15}$ ,  $a_{17}$ , and  $a_{23}$ . The central node is in the term with  $a_{14}$ . All of the other terms result from the cross-dispersive flux integrals of equations 3.1.1.8 and 3.1.1.9. Thus it will be advantageous to reduce the bandwidth of the final finite-difference equations by treating the cross-derivative dispersive-flux terms in an approximate manner. This will be covered in section 3.2.

Boundary cells with specified flux are handled similarly to interior cells. With the point-distributed grid, nodes will be located on boundary faces, edges, and corners. The cells associated with these boundary nodes will not have all eight subdomains (fig. 3.5). For example, a lateral boundary cell will have only four subdomains, while a corner boundary cell will have only one. The volume integrations over the cell are carried out as before, with the appropriate reduction in the number of subdomains. Fluxboundary conditions enter the finite-difference approximations through the surface integrations.

Consider a side node where part of the regional boundary is an x-plane; that is, the outward normal to the regional-boundary surface points in the positive x-direction. The associated half-cell for the node consists of four subdomains shown in figure 3.5. The discretization of equation 3.1.1.5 proceeds as follows. The temporal term for the rate-of-change-of-u becomes:

$$\frac{\partial}{\partial t} \int_{V_{m}} AudV \cong \frac{\partial}{\partial t} u_{i,j,k} \sum_{s=1}^{4} A_{ms} V_{ms}. \qquad (3.1.1.24)$$

The dispersive- and advective-flux terms are still given by equations 3.1.1.8 and 3.1.1.17, but now one of the faces, denoted by p=2 in figure 3.5, is a boundary face for the region. Note that its outward normal points in the positive x-direction. Using equation 3.1.1.1d for the flux-boundary condition, the integral over this face becomes:

$$\int_{S_{m2}} [\underline{\underline{B}} \cdot \nabla u - \underline{C}u] \cdot \underline{n} dS = - \int_{S_{m2}} \underline{J} \cdot \underline{n} dS ; \qquad (3.1.1.25a)$$

where

 $J_{xq}$  is the component of vector <u>J</u> in the x-direction in the qth element.

Normally  $J_x$  is constant over the entire cell face. Thus, the specified-flux boundary conditions has been incorporated in the finite-difference equation as a source term.

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The distributed-source and line-source terms simply are adjusted to account for the reduced cell volume. Thus equation 3.1.1.22 is unchanged, but equation 3.1.1.21b becomes:

 $\int_{V_m} DudV \cong u_{i,j,k} \sum_{s=1}^{\zeta} D_{ms} V_{s}. \qquad (3.1.1.26)$ 

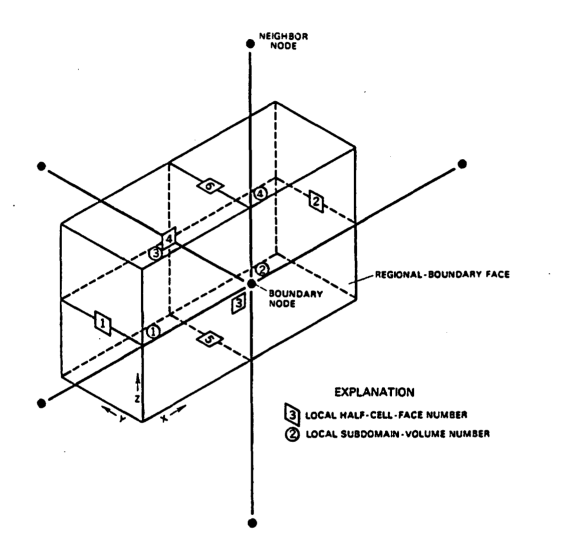


Figure 3.5.--Sketch of a boundary node with its half-cell volume showing the cell faces and the subdomain volumes.

In the flow equation, there is no advective term to cancel the corresponding term in the specified-flux boundary-condition equation. Inside the region, this means  $\underline{C} = 0$ ; but some of the boundary conditions have  $\underline{C} \neq 0$ . Then equation 3.1.1.9 takes the form:

$$\int_{S_{m2}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS = \int_{S_{m2}} [(\underline{\underline{C}} u) \cdot \underline{\underline{n}} - \underline{J} \cdot \underline{\underline{n}}] dS ; \qquad (3.1.1.27a)$$

which discretizes to:

$$\int_{S_{m2}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS \cong \sum_{q=1}^{4} [C_x u_{i,j,k} S_q - J_x S_q] . \quad (3.1.1.27b)$$

No central or upstream approximation for u is necessary, because the boundary face contains the node point.

# 3.1.2 Cylindrical Coordinates

The discretization of equation 3.1.1a in cylindrical coordinates is analogous to what has just been presented. We make the assumption of cylindrical symmetry, so no angular dependence exists. No line source terms • can be present so E=0. A cell volume becomes a ring bounded by  $r_{i-\frac{1}{2}}$  and  $r_{i+\frac{1}{2}}$ and  $z_{k-\frac{1}{2}}$  and  $z_{k+\frac{1}{2}}$ , where  $r_{i+\frac{1}{2}}$  is the radius of the cell wall between  $r_i$  and  $r_{i+\frac{1}{2}}$  (fig. 3.4).

An option is provided for automatic placement of the radial-grid lines between the interior and exterior radius. With this option the grid lines are spaced according to:

:

$$\frac{r_{i+1}}{r_i} = \left[\frac{r_{N_r}}{r_1}\right]^{1/(N_r-1)}$$

(3.1.2.1)

N<sub>r</sub> is the number of grid points (lines) in the radial direction;  $r_N_r$  is the exterior radius (m); and  $r_1$  is the interior radius (m).

where

This equation gives logarithmic-node spacing in the radial direction. The cell boundaries for the cylindrical faces are chosen to be at the logarithmic mean radii; for example:

$$\mathbf{r}_{i+\frac{1}{2}} = \frac{\mathbf{r}_{i+1} - \mathbf{r}_i}{\ln(\mathbf{r}_{i+1}/\mathbf{r}_i)}; \qquad (3.1.2.2)$$

for both the automatic and user-specified radial-grid distribution. A logarithmic-grid spacing will make the pressure drop uniform between adjacent grid points for steady radial flow in a homogenous medium (Aziz and Settari, 1979, p. 88). The discharge flux at  $r_{i+\frac{1}{2}}$  matches the analytical solution under these conditions.

Each cell ring is composed of four subdomain rings (shown in fig. 3.5). Thus, the temporal rate-of-change of u in the cell is approximated by:

$$\frac{\partial}{\partial t} \int_{V_{m}} AudV \cong \frac{\partial}{\partial t} u_{ik} \sum_{s=1}^{4} A_{ms} V_{ms}; \qquad (3.1.2.3)$$

where a sample subdomain volume is, for subdomain s=1:

$$V_1 = \pi \left( r_1^2 - r_{1-\frac{1}{2}}^2 \right) \frac{1}{2} \left( z_k - z_{k-1} \right) . \qquad (3.1.2.4)$$

The surface of cell m is composed for four faces, and each face belongs to two elements. Each element in the z-direction may have different porousmedium properties, but these properties must be constant in the r-direction. Equation 3.1.1.8 for the dispersive-flux term becomes:

$$\int_{S_{m}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS = \sum_{p=1}^{4} \sum_{q=1}^{2} \int_{S_{mpq}} (\underline{\underline{B}} \cdot \nabla u) \cdot \underline{\underline{n}} dS ; \qquad (3.1.2.5)$$

and typical integral over a face of a cell surface, p=2, is

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$$\int_{S_{m2}} (\underline{B} \cdot \nabla u) \cdot \underline{n} dS = \sum_{q=1}^{2} \int_{S_{m2q}} [B_{rr} \frac{\partial u}{\partial r} + B_{rz} \frac{\partial u}{\partial z}] dS ; \qquad (3.1.2.6a)$$

$$= \sum_{q=1}^{2} \left[ B_{rr}(q) \frac{\partial u}{\partial r} \right|_{i+\frac{1}{2},k} S_{m2q} + B_{rz}(q) \frac{\partial u}{\partial z} \left|_{q} S_{m2q} \right] \cdot (3.1.2.6b)$$

Equation 3.1.1.13, with r taking the place of x, is used to approximate  $\_$ 

 $\frac{\partial u}{\partial r}\Big|_{i+\frac{1}{2},k}$ , and equations 3.1.1.14 and 3.1.1.15 are used to approximate  $\frac{\partial u}{\partial z}\Big|_{q}$ .

Representative cell face areas are, for the face p=1:

$$S_{m_{11}} = 2\pi r_{i-\frac{1}{2}}(z_k - z_{k-1}); \qquad (3.1.2.7)$$

and, for the face p=3:

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$$S_{m_{21}} = \pi (r_1^2 - r_{1-\frac{1}{2}}^2);$$
 (3.1.2.8a)

$$S_{m_{32}} = \pi (r_{i+\frac{1}{2}}^2 - r_i^2) . \qquad (3.1.2.8b)$$

The advective term is treated as in the cartesian-coordinate case, with appropriate reduction in the number of faces and zones. Again, central or upstream differencing in space can be selected. The distributed source term proportional to u is also handled as in equations 3.1.1.21a and 3.1.1.21b with appropriate reduction in the number of subdomains from eight to four.

Finally, the finite-difference approximation to equation 3.1.1.1a for cylindrical coordinates with angular symmetry takes the form:

 $\frac{\partial}{\partial t} (a_{10} u_{i,k}) = a_1 u_{i-1,k-1} + a_2 u_{i,k-1} + a_3 u_{i+1,k-1}$ 

+ 
$$a_4 u_{i-1,k}$$
 +  $a_5 u_{i,k}$  +  $a_6 u_{i+1,k}$   
+  $a_7 u_{i-1,k+1}$  +  $a_8 u_{i,k+1}$  +  $a_9 u_{i+1,k+1}$  (3.1.2.9)

Note that no  $a_0$  term exists because annular-ring sources normally are not encountered. The terms  $a_2$ ,  $a_4$ ,  $a_6$ , and  $a_8$  are contributions from the closest neighbors to the central node point appearing in the  $a_3$  term. The other terms arise from the cross-derivative dispersive-flux integrals, and they may be treated in an approximate fashion to reduce the matrix-band width. The specified-flux boundary conditions are discretized in the same manner as for the cartesian-coordinate system, with appropriate reduction in the number of subdomains and surface faces.

#### 3.1.3 Temporal Discretization

To approximate the time derivative, two options are offered. The first is centered-in-time differencing, commonly known as the Crank-Nicholson method. The time derivative is approximated by the finite difference:

$$\frac{\partial}{\partial t} \left( a_{m} u_{m} \right) \cong \frac{\left( a^{m} u_{m} \right)^{n+1} - \left( a^{m} \right)^{n}}{t^{n+1} - t^{n}}; \qquad (3.1.3.1)$$

where t is the time at level n.

The right-hand-side, F, of the equation concerned is evaluated as follows:

$$F \cong \frac{1}{2}(F^{n+1} + F^{n});$$
 (3.1.3.2)

where

 $F^n$  is the spatial-difference function evaluated at time n.

The other option is backward-in-time differencing, which has the form:

$$\mathbf{F} \cong \mathbf{F}^{\mathbf{n+1}} \tag{3.1.3.3}$$

As with the advective spatial differencing, central-in-time differencing has the potential for causing oscillations in the solution (Price and others, 1966; Smith and others, 1977), whereas backwards-in-time differ-encing does not. However, backwards-in-time differencing does introduce numerical dispersion that must be kept under control by limiting the time-step size (Lantz, 1970; Price and others, 1966; Smith and others, 1977; Briggs and Dixon, 1968).

Equations 3.1.3.1 through 3.1.3.3 for the time discretization can be combined into a general form as:

$$\frac{(a_{m}u_{m})^{n+1} - (a_{m}u_{m})^{n}}{t^{n+1} - t^{n}} = \theta F^{n+1} + (1-\theta)F^{n}; \qquad (3.1.3.4)$$

where  $\theta = 1$  gives the fully implicit or backward-in-time (BT) differencing, and  $\theta = \frac{1}{2}$  gives the Crank-Nicholson or centered-in-time (CT) differencing.

The next step is to express the difference equation in residual form by writing:

$$u^{n+1} = u^n + \delta u$$
; (3.1.3.5)

where

ou is the temporal change in u.

Equation 3.1.3.5 is inserted in equation 3.1.3.4 and the following expansions of the temporal-difference terms are used. These are consistent differencing expansions that correspond to the differentials of products. For terms of the form  $(a_i u)^{n+1}$  we have:

$$(a_i u)^{n+1} - (a_i u)^n = a_i^{n+1} \delta u + u^n \delta a_i;$$
 (3.1.3.6)

for terms of the form  $(a_{i}a_{j}u)^{n+1}$ , we have:

$$(a_{i}a_{j}u)^{n+1} - (a_{i}a_{j}u)^{n} = a_{i}^{n+1}a_{j}^{n+1}\delta u + a_{i}^{n+1}u^{n}\delta a_{j} + a_{j}^{n}u^{n}\delta a_{i}.$$
 (3.1.3.7)

### 3.1.4 Finite-Difference Flow and Transport Equations

Combining equations 3.1.3.4, 3.1.3.6, and 3.1.3.7 with 3.1.1.23 or 3.1.2.9, we obtain the form of the general finite-difference equation for an interior node. The rather large number of terms makes presentation of the general equation impractical. It is more instructive to present the discretized flow, and heat and solute-transport equations individually, showing the x-direction terms only. The additional dispersive and convective terms for the y and z-directions follow the same pattern as their counterparts in the x-direction.

The finite-difference approximation to the flow equation (2.3.1a) is, for an interior node m:

$$C_{33}\delta p_{m} + C_{32}\delta T_{m} + C_{31}\delta w_{m} = \theta T_{Fi+\frac{1}{2}} (\delta p_{i+1} - \delta p_{i}) - \theta T_{Fi-\frac{1}{2}} (\delta p_{i} - \delta p_{i-1})$$

+ 
$$T_{F_{i+1}}(p_{i+1}^{n} - p_{i}^{n} + \rho_{i+1}^{n}g(x_{i+1} - x_{i}))$$

$$- \mathbf{r}_{Fi-\frac{1}{2}}(\mathbf{p}_{i}^{n} - \mathbf{p}_{i-1}^{n} + \boldsymbol{\rho}_{i-\frac{1}{2}}^{n}\mathbf{g} (\mathbf{x}_{i} - \mathbf{x}_{i-1}))$$
$$+ \mathbf{Q}_{m}^{n}\boldsymbol{\rho}^{\star} + \mathbf{\theta} \frac{\partial \mathbf{Q}_{m}^{n}}{\partial \mathbf{p}} \boldsymbol{\rho}^{\star} \delta \mathbf{p}_{i}$$

+ y and z direction difference terms; (3.1.4.1a)

where

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$$C_{33} = \left[\rho_{m}^{n+1} \sum_{s=1}^{8} \alpha_{bs} V_{s} + \rho_{o} \beta_{p} \sum_{s=1}^{8} \epsilon_{s}^{n} V_{ms}\right] / \delta t ; \qquad (3.1.4.1b)$$

$$C_{s2} = [\rho_0 \beta_T \sum_{s=1}^{8} \epsilon_s^n V_{ms}]/\delta t;$$
 (3.1.4.1c)

$$C_{31} = [\rho_0 \beta_W \sum_{s=1}^{8} \epsilon_s^n V_{ms}]/\delta t;$$
 (3.1.4.1d)

$$T_{Fi+\frac{1}{2}} = \frac{\rho_{i+\frac{1}{2}}^{n}}{\mu_{i+\frac{1}{2}}^{n}(x_{i+1}-x_{i})} \quad \begin{array}{c} 4 \\ \Sigma \\ q=1 \end{array} \quad k_{m2q} S_{m2q} ; \qquad (3.1.4.1e) \end{array}$$

$$T_{Fi-\frac{1}{2}} = \frac{\rho_{i-\frac{1}{2}}^{n}}{\mu_{i-\frac{1}{2}}^{n}(x_{i}-x_{i-1})} \xrightarrow{4}_{q=1} k_{mlq} S_{mlq}; \qquad (3.1.4.1f)$$

where

 $C_{ij}$  are the capacitance factors (various);  $T_{Fi}$  are the conductance terms for flow (m-s); and  $Q_m^n$  is the volumetric source flow rate for cell m (m<sup>3</sup>/s).

The flow-conductance factors,  $\tilde{T}_{Fi}$ , (m<sup>3</sup>), are defined as:

$$\tilde{T}_{Fi+\frac{1}{2}} = \sum_{q=1}^{4} k_{m2q} S_{m2q} / (x_{i+1} - x_i). \qquad (3.1.4.2)$$

These factors contain the spatial information and are constants.

In equation 3.1.4.1a, the source-sink flow rate has been made semi-implicit in time by including a term that accounts for changes in flow rate with changes in cell pressure. It is semi-implicit, because only the flow rate that contributes to the equation for cell m is treated implicitly. The effect of a change in pressure in cell m on the source-sink flow rates for other cells coupled to the cell through a well bore are not included; this approach avoids enlarging the bandwidth of the system equation matrix, equation 3.6.1a.

The finite-difference approximation to the hest-transport equation  
(2.3.1b) is, for an interior node, m:  

$$C_{23}6p_{m} + C_{22}6T_{m} + C_{21}6w_{m} = 6T_{Hi+\frac{1}{2}}(6T_{1+1}-6T_{1}) - 6T_{Hi-\frac{1}{2}}(6T_{1}-6T_{1-1}) + T_{Hi+\frac{1}{2}}(T_{1+1}^{n}-T_{1}^{n}) - T_{Hi-\frac{1}{2}}(T_{1}^{n}-T_{1-1}^{n}) - 6S_{xi+\frac{1}{2}}(T_{1}^{n}-T_{1-1}^{n}) - 6S_{xi+\frac{1}{2}}(T_{1}^{n}-T_{1-\frac{1}{2}}) - 6S_{xi+\frac{1}{2}}(T_{1-\frac{1}{2}}) - 7S_{xi+\frac{1}{2}}(T_{1-\frac{1}{2}}) - 7S_{xi+\frac{1}{2}}(T_{1-\frac{1}{2$$

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+ 
$$Q_m^n \rho^* c_f T_m^{*n} + \theta \frac{\partial Q_m^n}{\partial p} \rho^* \delta_m T_m^{*n} + \theta Q_m^n \rho^* c_f \delta_m^*$$

+ 
$$\theta \frac{\partial Q_m^n}{\partial p} \rho \star c_f \delta p_m \delta T_m^*$$

+ y and z direction dispersive, cross-dispersive and advective flux terms;

(3.1.4.3a)

where

$$C_{23} = \left[\rho_{0}\beta_{p}H_{m}^{n+1}\sum_{s=1}^{8} \varepsilon_{s}^{n}V_{s} + \rho_{m}^{n+1}H_{m}^{n+1}\sum_{s=1}^{8} \alpha_{s}V_{ms}\right]$$

$$-T_{m}^{n+1} \sum_{s=1}^{8} (\rho_{s}c_{s})_{s} \frac{\alpha_{bs}V_{ms}}{bs ms} / \delta t; \qquad (3.1.4.3b)$$

$$C_{22} = \left[\rho_{m}^{n+1} c_{f} \sum_{s=1}^{8} \varepsilon_{s}^{n} v_{s} + \sum_{s=1}^{8} (1-\varepsilon_{s}^{n})(\rho_{s}c_{s}) v_{s} + \sum_{s=1}^{8} (1-\varepsilon_{s}^{n})(\rho_{s}c_{s}) v_{s} + \frac{1}{2} (1-\varepsilon_{s}^{n})(\rho_{s}c_{s}) v_{s} + \frac{1$$

$$\rho_{o}\beta_{T}H_{m}^{n} \sum_{s=1}^{s} \sum_{ms}^{n} V_{ms} ]/\delta t ; \qquad (3.1.4.3c)$$

$$C_{21} = [\rho_0 \beta_W_m^n \sum_{s=1}^{8} v_{ms}]/\delta t;$$
 (3.1.4.3d)

$$T_{Hi+\frac{1}{2}} = \begin{bmatrix} \frac{4}{\Sigma} & \varepsilon_q^n & D_{Hxx}(2,q)S_{m2q} + K_f & \frac{4}{\Sigma} & \varepsilon_q^n & S_{m2q} + K_f & \frac{5}{Q-1} & \varepsilon_q^n & S_{m2q} \end{bmatrix}$$

$$\sum_{q=1}^{4} (1-\varepsilon_{q}^{n}) K_{sq} S_{m2q} ] / (x_{i+1} - x_{i}) ; \qquad (3.1.4.3e)$$

.

$$T_{Hi-\frac{1}{2}} = \begin{bmatrix} 4 \\ \Sigma \\ q=1 \end{bmatrix} = \begin{bmatrix} 4 \\ Q \\ Q \end{bmatrix} = \begin{bmatrix} 4 \\ Q \\$$

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$$\sum_{q=1}^{4} \frac{(1-\epsilon_q^n) K_{sq} S_{m1q}}{(1-\epsilon_q^n) K_{sq} S_{m1q}} ; \qquad (3.1.4.3f)$$

$$s_{xi+\frac{1}{2}}^{n} = (\rho v_{x})_{i+\frac{1}{2}}^{n} \frac{\xi}{q=1} \epsilon_{q}^{n} s_{m2q};$$
 (3.1.4.3g)

$$s_{xi-\frac{1}{2}}^{n} = (\rho v_{x})_{i-\frac{1}{2}}^{n} \frac{\xi}{q=1} \varepsilon_{q}^{n} S_{mlq};$$
 (3.1.4.3h)

$$\delta S_{xi+\frac{1}{2}} = \rho_{i+\frac{1}{2}}^{n} \delta v_{xi+\frac{1}{2}} \frac{4}{q=1} \epsilon_{q}^{n} S_{m2q} ; \qquad (3.1.4.3i)$$

$$\delta S_{xi-\frac{1}{2}} = \rho_{i-\frac{1}{2}}^{n} \delta v_{xi-\frac{1}{2}} \sum_{q=1}^{n} \varepsilon_{q}^{n} S_{nlq}; \qquad (3.1.4.3j)$$

$$T_{Hxy i+\frac{1}{2}} = \left[\sum_{q=1}^{4} \varepsilon_{q}^{n} D_{Hxy}(2,q) S_{m2q}\right] / (y_{j+1} - y_{j}); \qquad (3.1.4.3k)$$

$$T_{\text{Hxz } i+\frac{1}{2}} = \left[\sum_{q=1}^{4} \varepsilon_{q}^{n} D_{\text{Hxz}}(2,q) S_{m2q}\right] / (z_{k+1}^{-z} - z_{k}^{-z}); \qquad (3.1.4.32)$$

$$H = H(T_{oH}) + \overline{c_f} (T-T_{oH});$$
 (3.1.4.3m)

where  $T_{Hi}$  are the thermal conductance terms (W/°C).

In equation 3.1.4.3a, the same semi-implicit treatment of the source-sink flow rate has been incorporated as in equation 3.1.4.1a.

The central-or upstream-weighted value for the variables  $v_x$ ,  $\delta v_x$ , T and  $\delta T$  is given by the general form:

$$u_{i+k} = (1-\sigma) u_i + \sigma u_{i+1}$$
(3.1.4.4)

where

 $\sigma$  is the spatial weighting coefficient.

Central weighting is obtained with  $\sigma = \frac{1}{2}$ ; upstream weighting is obtained with  $\sigma = 0$  for a positive v.

The finite-difference approximation to the solute-transport equation (2.3.1c) is, for an interior node, m:

$$C_{13}\delta p_{m} + C_{12}\delta T_{m} + C_{11}\delta w_{m} = \theta T_{Si+\frac{1}{2}}(\delta w_{1+1} - \delta w_{1}) - \theta T_{Si-\frac{1}{2}}(\delta w_{1} - \delta w_{1-1})$$

$$+ T_{Si+\frac{1}{2}}(w_{1+1}^{n} - w_{1}^{n}) - T_{Si-\frac{1}{2}}(w_{1}^{n} - w_{1-1}^{n})$$

$$-\theta S_{xi+\frac{1}{2}}^{n+1} \delta w_{1+\frac{1}{2}} - \theta \delta S_{xi+\frac{1}{2}}w_{1+\frac{1}{2}}^{n}$$

$$+ \theta S_{xi+\frac{1}{2}}^{n+1} \delta w_{1-\frac{1}{2}} + \theta \delta S_{xi-\frac{1}{2}}w_{1-\frac{1}{2}}^{n}$$

$$- S_{xi+\frac{1}{2}}^{n}w_{1+\frac{1}{2}} + S_{xi-\frac{1}{2}}^{n}w_{1-\frac{1}{2}}^{n}$$

$$+ T_{Sxy i+\frac{1}{2}}(w_{1+1,j+1,k}^{n} + w_{1,j+1,k}^{n} - w_{1+1,j+1,k}^{n} - w_{1,j-1,k}^{n})$$

$$+ T_{5xz i+\frac{1}{2}}(w_{1+1,j+1,k}^{n} + w_{1,j+1,k}^{n} - w_{1+1,j+1,k}^{n} - w_{1,j,k-1}^{n})$$

$$- T_{5xy i-\frac{1}{2}}(w_{1,j+1,k}^{n} + w_{1-1,j+1,k}^{n} - w_{1,j-1,k}^{n} - w_{1-1,j-1,k}^{n})$$

$$- T_{5xz i-\frac{1}{2}}(w_{1,j,k+1}^{n} + w_{1-1,j,k+1}^{n} - w_{1,j,k-1}^{n} - w_{1-1,j,k-1}^{n})$$

$$- \lambda M_{m}^{m} w_{m}^{n} - \theta \lambda (w_{m}^{n+1} \delta M_{m} + M_{m}^{m} \delta w_{m})$$

$$+ 0_{m}^{m} \rho^{\alpha} w_{m}^{m} + \theta \frac{\partial Q_{m}^{m}}{\partial p_{p}} \delta p_{m} \rho^{\alpha} w_{m}^{m} + \theta Q_{m}^{m} \delta (\rho^{\alpha} w^{\alpha})_{m}$$

$$+ \theta \frac{\partial Q_{m}^{m}}{\partial p_{p}} \delta p_{m} \delta (\rho^{\alpha} w^{\alpha})_{m}$$

$$+ y \text{ and } z \text{ direction dispersive, cross-dispersive and advective-flux terms;} (3.1.4.5a)$$

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where

$$C_{13} = \left[\rho_{0}\beta_{p} w_{m}^{n} \sum_{s=1}^{8} K_{s}^{n} V_{s} + \rho_{m}^{n+1} w_{m}^{n+1} \sum_{s=1}^{8} \alpha_{bs} V_{ms}\right] / \delta t \qquad (3.1.4.5b)$$

$$C_{12} = [\rho_0 \beta_T w_m^n \sum_{s=1}^{\delta} k_s^n v_{ms}]/\delta t ; \qquad (3.1.4.5c)$$

$$C_{11} = [\rho_0 \beta_w w_m^n + \rho_m^{n+1}] \frac{g_k K_s^n V_m}{s=1} \sqrt{\delta t}; \qquad (3.1.4.5d)$$

$$\tau_{\text{Si+l_2}} = \begin{bmatrix} 4 \\ \Sigma \\ q=1 \end{bmatrix} \begin{bmatrix} 2 \\ q \end{bmatrix} \begin{bmatrix} 4 \\ Q \\ Q \end{bmatrix} \begin{bmatrix} 2 \\ Q \\ Q \end{bmatrix} \end{bmatrix} \begin{bmatrix} 2 \\ Q \\ Q \end{bmatrix} \begin{bmatrix} 2 \\ Q \\ Q \end{bmatrix} \end{bmatrix} \begin{bmatrix} 2 \\ Q \\ Q \end{bmatrix} \begin{bmatrix} 2 \\ Q \\ Q \end{bmatrix} \end{bmatrix} \end{bmatrix} \begin{bmatrix}$$

$$T_{\text{Si-l}_{2}} = \left[\sum_{q=1}^{4} \varepsilon_{q}^{n} D_{\text{Sxx}}(1,q) S_{mlq} + D_{m} \sum_{q=1}^{4} \varepsilon_{q}^{n} S_{mlq}\right] \frac{\rho_{i-l_{3}}^{n}}{x_{i}-x_{i-i}}; \quad (3.1.4.5f)$$

$$T_{\text{Sxy } i+\frac{1}{2}} = \left\{ \sum_{q=1}^{4} \varepsilon_{q}^{n} \quad D_{\text{Sxy}}(2,q) S_{m2q} \right\} \quad \frac{\rho_{i+\frac{1}{2}}}{y_{j+1} - y_{j}} ; \quad (3.1.4.5g)$$

$$T_{Sxz \ i+\frac{1}{2}} = \begin{bmatrix} 4 \\ 2 \\ q=1 \end{bmatrix} p_{Sxz}^{n} (2,q)S_{m2q} = \begin{bmatrix} \rho_{i+\frac{1}{2}}^{n} \\ z_{k+1} \end{bmatrix} (3.1.4.5h)$$

$$\mathcal{M}_{m}^{n} = \rho_{m}^{n} \sum_{s=1}^{8} \mathcal{K}_{s}^{n} V_{s} ; \qquad (3.1.4.5i)$$
$$\delta \mathcal{M}_{m} = \rho_{m}^{n+1} \sum_{s=1}^{8} \alpha_{bs} V_{s} \delta p_{m}$$

$$+ \sum_{s=1}^{\delta} (\kappa_s^n V_s) [\rho_o \beta_p \delta p_m + \rho_o \beta_T \delta T_m + \rho_o \beta_w \delta w_m]; \qquad (3.1.4.5j)$$

$$\mathbb{K}_{s}^{n} = \varepsilon_{s}^{n} + (\rho_{b}K_{d})_{s};$$
 (3.1.4.5k)

where

M<sup>n</sup><sub>m</sub> is the mass of fluid plus the effective additional fluid mass from sorption in cell m at time level n (kg); T<sub>Si</sub> are the conductance terms for solute transport (kg/s); and K<sub>s</sub> is the augmented porosity factor for subdomain s(-). In equation 3.1.4.5a, the same semi-implicit treatment of the source-sink flow rate has been incorporated as in equation 3.1.4.1a. In equations 3.1.4.1a-f, 3.1.4.3a-L, and 3.1.4.5a-k, subscripts pertaining to the y and z directions have been omitted for clarity, unless necessary. The source density,  $\rho^*$ , temperature, T\*, and mass fraction, w\*, are specified functions of time and source location. When the source-flow rate is negative, so that it becomes a sink, the density, temperature, and mass fraction become those of the cell. In the abbreviated subscript notation,  $u_m$  and  $u_i$  become identical for a given variable, u. Note that the cross-dispersive flux terms have been evaluated explicitly, that is, at time n, to limit the number of elements in the coefficient matrix of the unknowns,  $\underline{A}$ , to a maximum of seven for each equation. The coefficients  $T_i$ ,  $S_i$ , and  $N_i$  are evaluated at time n.

The preceding flow and transport equations are valid for confined flow. The forms of the capacitance terms that contain the porous-medium bulk compressibility are based on a slightly compressible porous matrix and a cell volume that deforms slightly in space. The coefficients  $C_{ij}$ , and the cell facial areas  $S_{mpq}$  are modified for the case of unconfined flow, as will be shown in section 3.4.6.

The permeability tensor in the flow equation is a diagonal matrix in the numerical implementation, because the coordinate directions are chosen to be along the principal directions of this tensor. These directions are assumed not to change with position in the simulation region. The finite-element discretization technique must be used for the more general situation of spatially variable, anisotropic, permeability directions.

In summary, the properties and variables that are spatially discretized on a cell-by-cell basis include pressure, temperature, solute-mass fraction, density, viscosity, enthalpy, and specified fluxes. Porous-matrix properties that are discretized on an element-by-element or zonal basis include porosity, permeability, thermal conductivity, heat capacity, bulk compressibility, bulk density, equilibrium-distribution coefficient, longitudinal dispersivity, and transverse dispersivity. Well-completion intervals also are designated on a zonal basis.

#### 3.1.5 Numerical Dispersion and Oscillation Criteria

For guidance in selecting the spatial and temporal discretization method, the following results have been obtained by Lantz (1970), Roache (1976, p. 19, 48), Smith and others (1977) and Price and others (1966), expressing the truncation errors that give rise to numerical dispersion and criteria for avoiding oscillations in the solution. They were derived for the onedimensional form of equation 3.1.1a with constant coefficients and no source terms; that is:

$$A \frac{\partial u}{\partial t} = B \frac{\partial^2 u}{\partial x^2} - C \frac{\partial u}{\partial x} . \qquad (3.1.5.1)$$

Similar analyses can be performed for the more general equation:

$$\frac{\partial u}{\partial t} = \frac{\partial}{\partial x} \left( B \frac{\partial u}{\partial x} \right) - C \frac{\partial u}{\partial x} + D u + E ; \qquad (3.1.5.2)$$

where B, C, D, and E are functions of x and t, and A is positive.

The truncation errors and oscillation criteria for both equation forms are given in table 3.1. The maximum values of A, B, and C should be used in the variable coefficient case, equation 3.1.5.2. All of the methods are stable in the sense that errors do not grow without bound. However, oscillations in space and time may persist without growth or decay. The oscillation criterion for the centered-in-time differencing was presented by Keller (1960, p. 140) and Briggs and Dixon (1968). They are sufficient conditions; thus, they may be conservative. Alternate conditions appear in Price and others (1966) but they require knowledge of the maximum or minimum eigenvalue of the spatial-discretization matrix that cannot be expressed analytically. An important thing to note from table 3.1 is that it is possible for oscillations in the solution to arise from both spatial and temporal discretization. For the flow equation with no advective term, oscillations from temporal discretization are still possible. For the flow equation in

cylindrical coordinates, an advective-type term appears so oscillations can also be caused by spatial discretization. If a source term that depends on u appears in equation 3.1.5.2, the oscillation criteria for the centered-in-time discretization are modified as shown.

When using the backwards-in-space (upstream) or backward-in-time differencing, one needs to check that the truncation-error terms that cause numerical dispersion do not become large relative to the physical-dispersion coefficient. Mathematically, for a dispersion coefficient given by equation 2.2.6.1.2; one needs to adhere to the following criteria:

$$\frac{\Delta x}{2} << \alpha_{\rm L}$$
; (3.1.5.3)

(3.1.5.4)

and

$$\frac{C\delta t}{2} \ll \alpha_{L}$$
;

where

$$\Delta x = x_{i+1} - x_i$$
; and (3.1.5.5a)

$$\delta t = t^{n+1} - t^n . (3.1.5.5b)$$

Note that these results are from a one-dimensional analysis with constant coefficients, but they give guidance for grid and time-step selection. Table 3.1 shows that, in the case of variable coefficients, additional truncationerror terms occur with backwards-in-time differencing, that can give rise to numerical-dispersion errors.

# Table 3.1.--Truncation errors and oscillation criteria for onedimensional parabolic equations

[BS, backward-in-space; BT, backward-in-time; CS, centered-in-space; CT, centered-in-time;

Discretization Method	Truncation Error	Oscillation Criterion
Equation 3.1.5.1		— <u> </u>
BS	$\frac{C\Delta x}{2}uxx$	
BT	$\frac{C^2 \Delta t}{2} \mathbf{u}_{\mathbf{x}\mathbf{x}}$	
CS	0(Δx <sup>2</sup> )	$\Delta x \leq \frac{2B}{ C }$
CT	0(Δt²)	$\frac{\Delta t}{(\Delta x)^2} \leq \frac{\lambda}{B}$

$$u_{xx} = \frac{\partial^2 u}{\partial x^2}$$
;  $u_x = \frac{\partial u}{\partial x}$ .]

Equation 3.1.5.2

BS

BT

$$\frac{C\Delta x u}{2} x x - \frac{C^2 \Delta t u}{2} x x + B_t \Delta t u x + B D u x x + 3B_x C u x x + 2B_x^2 u x x - \frac{1}{2} x + 3B_x C u x x + 2B_x^2 u x x + 2B_x^2 u x x + \frac{1}{2} x + \frac{1}{2$$

CS 
$$0(\Delta x^2)$$
  $\Delta x \leq \frac{2B}{|C|}$   
CT  $0(\Delta t^2)$   $\Delta t \leq MIN \left(\frac{1}{B} - \frac{D}{2}, \frac{2}{D}\right); D>0$ 

. .

$$\Delta t \leq \frac{1}{\frac{B}{\Delta x^2} \cdot \frac{D}{2}}; D \leq 0$$

3.1.6 Automatic Time-Step Algorithm

Manual time-step selection can be difficult, when many source terms and boundary conditions change considerably with time. In general, the more rapidly the conditions change, the smaller the time steps will need to be for an accurate solution. Therefore, the heat- and solute-transport simulator has an automatic time-step option that uses an empirical algorithm (INTERCOMP Resource Development and Engineering, Inc., 1976). The user specifies the maximum values of change in pressure, temperature, and mass fraction considered acceptable as well as the maximum and minimum time step allowed. Then, at the beginning of each time step, the following adjustments are made, depending on the conditions:

if: 
$$|\delta u_{\max}| > \delta u_{\max}^{s}$$
;  $\delta t = \frac{1}{2} \delta t_{o} (1 + \frac{\delta u_{\max}^{s}}{|\delta u_{\max}|})$ ; (3.1.4.1)

otherwise, if:

$$0 < |\delta u_{max}| < \delta u_{max}^{s}; \delta t = \delta t_{o}(0.2 + 0.8 \frac{\delta u_{max}^{s}}{|\delta u_{max}|});$$
 (3.1.4.2)

otherwise, if:

$$\delta u_{max} = 0; \ \delta t = 1.5 \delta t_{o};$$
 (3.1.4.3)

where

u is pressure, temperature, or mass fraction;

 $\delta u_{max}^{S}$  is the specified maximum change in u;

St is the new time step;

St is the previous time step; and

 $\delta u_{max}$  is the absolute value of the maximum-calculated change in u over the

previous time step.

The new time step is selected to be the minimum of the three that were calculated on the basis of change in the pressure, temperature, and mass fraction. The time step is constrained to a user-specified range, and the maximum increase in ôt is limited to a factor of 1.5. This algorithm tends to increase the time step such that the maximum acceptable change in pressure, temperature, or mass fraction is achieved as the simulation progresses. The minimum required time step, set by the user, is maintained for the first two steps after boundary-condition changes occur or after the automatic time-step algorithm is invoked.

# 3.1.7 Discretization Guidelines

No complete set of discretization rules exists that will guarantee an accurate solution discretization with a minimum number of nodes and time steps, even for the case of constant coefficients. However, the following empirical guidelines should be useful.

- If using the backward-in-space or backward-in-time differencing, make some estimates of the truncation error, using parameter values at their limits expected for the simulation. Thus, verify that the grid-spacing and time-step selection do not introduce excessive numerical dispersion.
- 2. If using centered-in-space and centered-in-time differencing, print results every time step for a short simulation period, 5-10 time steps. Examine the results for spatial and temporal oscillations that are caused by the time or space discretization being too coarse.
- 3. Check on spatial-discretization error by refining the mesh. However, this often is impractical for large regions. A check on temporal-discretization error is relatively easy to make by refining the time-step length for a short simulation.

4. At each change of boundary condition or source flow rates, reduce the time step until the abrupt changes have had time to propagate into the region. The automatic time-step algorithm does this.

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- 5. To adequately represent a sharp solute-concentration or temperature front, span it with at least 4-5 nodes. A large number of nodes may be required if a sharp front moves through much of the region over the simulation time. Compromises often will have to be made. An advantage of the centered-in-space differencing is that oscillations will reveal when the grid is too coarse relative to the gradients of solute concentration or temperature.
- 6. Well flows that highly stress the aquifer require a small time step after a change in flow rate, to control errors from explicit flow-rate allocation or explicit well-datum pressure calculation.
- 7. Sometimes, the global-balance summary table will indicate that the time step is too large by exhibiting large residuals, particularly if the density and viscosity variations are large.
- To check for unusual results that could indicate discretization error, print out all of the results some of the time, and some of the results all of the time.

#### 3.2 PROPERTY FUNCTIONS AND TRANSPORT COEFFICIENTS

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Numerical implementation of the fluid-density function is simply the evaluation of equation 2.2.1.1b or 2.2.1.3a. Fluid viscosity is obtained by evaluation of equation 2.2.2.1 and equation 2.2.2.2 if necessary. The enthalpy of pure water at the selected reference values of pressure and temperature,  $H(p_{OH}, T_{OH}, 0)$  is obtained by a two-step interpolation. First, the enthalpy of saturated fluid at the given temperature is calculated by linear interpolation in the table of saturated enthalpy as a function of temperature; then, adjustment to the given pressure is made by bilinear interpolation in

the table of enthalpy deviation from saturation as a function of pressure and temperature. This procedure is given by equation 2.2.3.1a. A sequential search is made for each interpolation, since the number of pressure or temperature entries is 32 or less in both tables. Equation 2.2.3.1c is used for all subsequent fluid-enthalpy calculations. It is possible that simulation of wide variations in pressure and temperature could require a table look-up for all enthalpy calculations, and the algorithm in the program code would need to be modified.

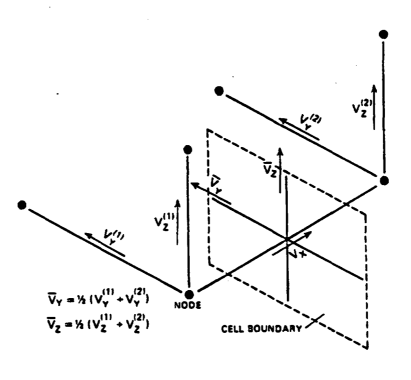


Figure 3.6.--Sketch of velocity vectors used for the dispersioncoefficient calculation for a given cell. The hydrodynamic-dispersion coefficient is calculated by equation 2.2.6.1.1 with equations 2.2.6.1.2 and 2.2.6.1.3. A separate value of  $D_{sij}(p,q)$  is associated with each element, q, of each cell face, p. Interstitial velocities are obtained from the pressure and elevation differences across the face for the velocities normal to the face. Velocities parallel to the face are determined by averaging velocities from each side of the face. An x-face, with the y and z velocities interpolated to get the effective values on the subface appears in figure 3.6. Average values are used, since the face lies midway between  $x_i$  and  $x_{i+1}$ .

The thermo-hydrodynamic-dispersion tensor is calculated by equations 2.2.6.2.1 and 2.2.6.2.2. The porosity and the thermal conductivities are defined by zones and the interstitial velocities are obtained the same as for the hydrodynamic dispersion.

Two methods are available in the HST3D simulator for computation of the cross-derivative dispersive-flux terms. The most rigorous treatment of the cross-derivative terms involves explicit calculation. They are lagged one iteration in the solution cycle of the flow, heat, and solute equations. The cross-derivative dispersive fluxes are recalculated for each iteration based on the conditions existing at the end of the previous iteration and then they are incorporated into the right-hand-side vector. Therefore at least two iterations in the solution cycle are required at each time step. This full treatment requires storage of the nine dispersion-coefficient terms for thermal and solute dispersion. An approximate empirical treatment of the cross-derivative dispersion terms is available also, that consists of lumping the cross-derivative dispersion coefficients into the diagonal dispersioncoefficient terms. The three augmented dispersion coefficients for thermal and solute dispersion are the only coefficients stored, and extra iterations are not required, because the cross-derivative dispersive fluxes are not computed.

#### 3.3 SOURCE OR SINK TERMS--THE WELL MODEL

In the present version of the HST3D program, only one well can exist in a particular cell. Multiple wells in a cell must be represented by an equivalent single well, or the spatial grid must be refined to separate them. This restriction includes wells that are located in the same areal cell that are completed in different vertical intervals.

Recall that a cell may contain up to four zones of different porous-media properties over a given areal plane. If a well is completed in a cell with multiple zones, the effective ambient permeability is taken to be that of the lowest zone number. This is because no algorithm presently exists to calculate the effective ambient permeability for a well in areally heterogeneous porous media.

## 3.3.1 The Well-Bore Model

The volumetric flow rate per unit length of well bore is given by equation 2.4.1.1. Discretization for a given cell, m, is achieved by choosing the average pressure to be the cell pressure, and multiplying by the length of well bore in that cell. Since the well bore is usually screened over the more permeable zones of the formation region, the screened intervals are specified by zones or sets of elements rather than by cells. The upper and lower parts of a screened interval will be one-half of the cell thickness in length, unless the cell in question is an upper or a lower boundary cell for the region. Thus:

$$Q_{wl} = \frac{(p_{wl} - p_m) [W_I(l_1)L_{l_1} + W_I(l_2)L_{l_2}]}{\mu_m(l)}$$
(3.3.1.1a)

In the case of an unconfined aquifer with a well screened through the free surface, the screened length,  $L_g$ , is adjusted as the saturated thickness varies in time.

where

$$L_{g_1} = \frac{1}{2}(z_g - z_{g_1}) ; \qquad (3.3.1.1b)$$

$$L_{g2} = \frac{1}{2} (z_{g+1} - z_g) ; \qquad (3.3.1.1c)$$

and where

 $Q_{w\ell}$  is the volumetric-flow rate from the well to the aquifer in cell m at well-bore level  $\ell$  (m<sup>3</sup>/s);

p<sub>m</sub> is the pressure at node m (Pa);

- $p_{wl}$  is the pressure in the well bore at elevation of node m (Pa);
- m(l) is the cell number associated with the well-bore level l;
- $L_{\ell_1}$  is the length of well bore in the lower half of cell m( $\ell$ ) (m); and

 $L_{g_2}$  is the length of well bore in the upper half of cell m( $\ell$ ) (m).

Equations 3.3.1.1b and 3.3.1.1c are valid for the z-coordinate directed vertically upward.

For wells drilled at an angle  $\theta_{ij}$  to the vertical:

L<sub>2</sub> =

$$L_{\ell 1} = \frac{\frac{1}{2} (z_{\ell} - z_{\ell-1})}{\cos \theta_{u}}; \qquad (3.3.1.2a)$$

(3.3.1.2b)

The well indices may be different in the upper and lower halves of the cell, because the porous-medium zone boundaries pass through planes of node points. The two-term sum in equation 3.3.1.1a accounts for this. When the cell is at the upper or lower boundary of the region, or at the ends of the screened interval for the well, the appropriate term in equation 3.3.1.1a becomes zero.

For notational convenience, we define:

$$H_{wl}L_{l} = M_{wl1}L_{l1} + M_{wl2}L_{l2}; \qquad (3.3.1.3)$$

where  $M_{c}$  is the well mobility defined by equation 2.4.1.5.

Flow-rate allocation by mobility is obtained by discretizing equation 2.4.1.5 to give:

$$Q_{w\ell} = \frac{Q_w H_{w\ell} L_{\ell}}{\sum_{\ell=\hat{\ell}_L}^{U} H_{w\ell} L_{\ell}}; \qquad (3.3.1.4)$$

where

 $\hat{L}_L$  is the index of the bottom level of the well screen; and  $\hat{L}_R$  is the index of the top level of the well screen.

If the screened interval is not continuous from  $l_L$  to  $l_U$ , the length,  $L_{lj}$  is set to zero over the appropriate subintervals. For an observation well, the dependent variable data in the aquifer are taken from the cell at location  $l_U$ . Flow-rate allocation by the product of mobility and pressure difference is obtained by discretization of equations 2.4.1.9 using equation 2.4.1.8. The pressure at the well datum is given by (Thomas, 1982, p. 155):

$$P_{wd} = \frac{\sum_{\ell=l_{L}}^{l} M_{w\ell} L_{\ell}(P_{m} + P_{w\ell} z_{\ell}) - Q_{w}}{\sum_{\ell=l_{L}}^{l} M_{w\ell} L_{\ell}};$$

(3.3.1.5)

and the flow rate from the well to the aquifer at each layer is given by:

$$Q_{w\ell} = M_{w\ell} L_{\ell} [p_{wd} + \rho_{w} g(z_{wd} - z_{\ell}) - p_{m}];$$
 (3.3.1.6)

where  $z_{g}$  is the elevation of the well node at level  $\mathcal{L}$  (m).

Similar expressions were derived by Bennett and others (1982) for constant-density fluids.

For simulations with a well completed in more than one layer, and explicit calculation of the well-datum pressure, a large well index or mobility can cause computational instabilities (Chapplear and Williamson, 1981). A large flow rate will be allocated to a layer with large mobility, and the cell pressure can become nearly equal to the well-bore pressure. This will make the flow-rate allocation small during the next time step, and an oscillation may develop. To avoid a severe time-step limitation, a semiimplicit, well flow-rate allocation can be used. It is available as a calculation option in the HST3D program. Equation 3.3.1.6 becomes:

 $Q_{w\ell}^{n+1} = H_{w\ell}L_{\ell}(p_{wd}^{n} + \rho_{w}^{n} g(z_{wd}^{-}z_{\ell}) - p_{m}^{n}) - H_{w\ell}L_{\ell}\delta p_{m} . \qquad (3.3.1.7)$ 

This gives an implicit coefficient that is included in the matrix element for node m in the finite-difference equations. Note that well-datum pressure still is treated explicitly. This can put a restriction on the time step for stability particularly when the aquifer is being stressed heavily. Also the total flow rate in the well will not be maintained over the time step. Therefore, iterations are necessary.

A fully implicit approach would eliminate the iterations, but would introduce additional coefficients in the flow equation for all the cells that were communicating with the given well. The band-width of the finitedifference flow equations, would be increased, thus making the two matrixsolution techniques, much more difficult to implement. However, Bennett and others (1982) employ the fully implicit approach with a compatible matrixsolution technique.

A compromise algorithm was developed starting from equation 3.3.1.5 expressed as a well constraint to maintain specified well-flow rate:

$$\sum_{l=l_{L}}^{l} M_{wl} L_{l} \delta p_{m} - \sum_{l=l_{L}}^{l} M_{wl} L_{l} \delta p_{wd} = 0. \qquad (3.3.1.8)$$

Equation 3.3.1.8 is written for each well in the region.

The matrix representation of the flow- and well-constraint equations being solved simultaneously is bordered as shown by:

 $\begin{bmatrix} \underline{A} & \underline{W}_{2} \\ \underline{W}_{1} & \underline{W}_{3} \end{bmatrix} \begin{bmatrix} \underline{b}_{1} \\ \underline{0}_{2} \\ \underline{0}_{2} \end{bmatrix} = \begin{bmatrix} \underline{b}_{1} \\ \underline{0} \end{bmatrix}; \qquad (3.3.1.9)$ 

where

- $\underline{A}$  is the coefficient matrix from the discretized flow equation;  $\underline{W}_1$  is the coefficient matrix linking the pressures in each of
- the cells which communicate with a well;
- $\underline{W}_2$  is the coefficient matrix linking the well-datum pressures to the flow equation through the source term;

- $\underline{W}_3$  is the coefficient matrix (diagonal) for the well-datum pressures in the well constraint equation (eq. 3.3.1.8); and
- $\underline{b}_1$  is the vector of known quantities from the discretized flow equation.

The two equations are solved iteratively at each time step for  $\underline{\delta p}$  and  $\underline{\delta p}_{wd}$ . The well-datum pressures are lagged one iteration in the solution of the flow equations. The initial value for  $\underline{\delta p}_{wd}$  is taken to be  $\underline{0}$ . The iterations are terminated when the maximum fractional change in  $\underline{\delta p}_{wd}$  is less than 0.001. Usually, only two or three iterations are required for convergence. This algorithm has the advantage that the sparse structure of the matrix  $\underline{A}$  is preserved, so that the implemented matrix-equation solvers can be employed. At the first time step, equation 3.3.1.4 is used to calculate the flow rates at each layer for each well. Equation 3.3.1.6 is used thereafter.

A reversal of flow between the well and the aquifer at any layer communicating with a well is allowed. However, difficulties arise if there is a reversal of flow within the well bore. An algorithm to compute a realistic density profile in the well bore under flow-reversal conditions has not yet been developed; therefore, the following algorithm is currently used in HST3D to compute heat and solute flow rates in a well bore.

For a production well, heat and solute balance calculations are done from the bottom to the top of the well-screen interval. If injection occurs at a given layer, the density, temperature, and solute concentration injected are based on the current values coming up the well bore from below. Any fluid flowing down the well bore to that layer is neglected. Density, temperature, and solute concentration values based on well-datum conditions are used if there is no upward flow in the well bore below the given injection layer. This algorithm is suitable for producing wells which leak into the aquifer but have net upward flow along the entire well bore. It may be a poor approximation if there are large density, temperature, or solute concentration variations within the well and flow reversals occur in the well bore.

For an injection well, no account is taken of the effect of producing layers on the density, temperature, or solute concentration in the well bore. The conditions at the well-datum level are used for all injection layers. Clearly, this approximation is valid only for injection wells with slight invasion from producing layers and no flow reversals in the well bore. A more realistic algorithm will require a complex, iterative calculation.

In the present version of the HST3D simulator, when a production or injection well becomes inactive, by having its flow rate set to zero, no circulation of fluid from one aquifer-discretization layer to another is computed. Removing this restriction would require the algorithm, described previously, to handle flow reversals in the well bore.

For the case of a single well in the cylindrical-coordinate system, equation 2.4.1.12, for the well bore, is discretized in space and time in the same manner as the system flow equation. Flow-rate allocation by mobility and pressure gradient or specified pressure at the well datum are the options available. The augmented-flow equation 2.4.1.13, is discretized in space and time in the manner that led to equation 3.1.4.1a. At a node along the well screen below the top of the screen,  $k < l_{11}$ , the equation is:

$$C_{33}\delta p_{m} + C_{32}\delta T_{m} + C_{31}\delta w_{m} = \theta(T_{Fk+\frac{1}{3}} + T_{wFk+\frac{1}{3}})(\delta p_{k+1} - \delta p_{k})$$
  
- $\theta(T_{Fk-\frac{1}{3}} + T_{wFk-\frac{1}{3}})(\delta p_{k} - \delta p_{k-1})$   
+ $(T_{Fk+\frac{1}{3}} + T_{wFk+\frac{1}{3}})(p_{k+1}^{n} - p_{k}^{n}) + T_{Fk+\frac{1}{3}}\rho_{k+\frac{1}{3}}^{n}g(z_{k+1} - z_{k})$   
+ $T_{wFk+\frac{1}{3}}\rho_{wk+\frac{1}{3}}^{n}g(z_{k+1} - z_{k})$   
- $(T_{Fk-\frac{1}{3}} + T_{wFk-\frac{1}{3}})(p_{k}^{n} - p_{k-1}^{n}) - T_{Fk-\frac{1}{3}}\rho_{k-\frac{1}{3}}^{n}g(z_{k} - z_{k-1})$   
- $T_{wFk-\frac{1}{3}}\rho_{wk-\frac{1}{3}}^{n}g(z_{k} - z_{k-1})$ 

+ r-direction dispersive terms; (3.3.1.10a)

where

$$T_{wFk+\frac{1}{2}} = \frac{4\pi r_w^3}{v_w f_w (z_{k+1} - z_k)};$$

(3.3.1.10b)

 $T_{wFk-\frac{1}{2}} = \frac{4\pi r_w^3}{v_w f_w (z_k - z_{k-1})};$ 

(3.3.1.10c)

and where

 $T_{\rm up}$  are the conductances for flow at the well bore (m-s).

For the node at the top of the well-screen interval,  $k=\lambda_U$ , the discretized, augmented flow equation (2.4.1.13) becomes:

$$C_{33}\delta p_{m} + C_{32}\delta T_{m} + C_{31}\delta w_{m} = -\theta(T_{F\bar{P}_{U}^{-\frac{1}{2}}} + T_{wF\bar{P}_{U}^{-\frac{1}{2}}})(\delta p_{\bar{P}_{U}} - \delta p_{\bar{P}_{U}^{-1}})$$
$$-(T_{F\bar{P}_{U}^{-\frac{1}{2}}} + T_{wF\bar{P}_{U}^{-\frac{1}{2}}})(p_{\bar{P}_{U}}^{n} - p_{\bar{P}_{U}^{-1}}^{n}) - T_{F\bar{P}_{U}^{-\frac{1}{2}}}p_{\bar{P}_{U}^{-\frac{1}{2}}}^{n}p_{\bar{P}_{U}^{-\frac{1}{2}$$

+ r-direction dispersive terms;

(3.3.1.11)

where

 $Q_w$  is the specified volumetric flow rate of the well (positive is injection to the aquifer) (m<sup>3</sup>/s).

In the case of specified pressure at the well datum, equation 3.3.1.11 is replaced by:

Pl, = Pwd

(3.3.1.12)

The well-bore velocity and friction factor are calculated explicitly at the beginning of the time step. Since the friction factor is a weak function of velocity, this causes no instabilities. Evaluating the well-conductance factors explicitly is consistent with the treatment of the aquifer-conductance factors.

#### 3.3.2 The Well-Riser Model

The well-riser calculation is done by numerically solving equation 2.4.2.11. These ordinary differential equations are integrated using the midpoint method with rational-function extrapolation, developed by Bulirsch and Stoer (1966) and presented by Gear (1971, p. 96).

The following algorithm is applied to the well-riser calculations:

$$p_r^{\star} = p_{rk} + \frac{\Delta \ell}{2} F(p_{rk}, T_{rk}, \ell_k);$$
 (3.3.2.1a)

$$T_r^* = T_{rk} + \frac{\Delta \ell}{2} G(p_{rk}, T_{rk}, \ell_k);$$
 (3.3.2.1b)

$$p_{rk+1} = p_{rk} + \Delta \ell F(p_r^*, T_r^*, \ell_k + \frac{\Delta \ell}{2});$$
 (3.3.2.1c)

$$T_{rk+1} = T_{rk} + \Delta l G(p_r^*, T_r^*, l_k + \frac{\Delta l}{2});$$
 (3.3.2.1d)

where

$$\Delta \mathbf{r} = \mathbf{r}_{k+1} - \mathbf{r}_k. \tag{3.3.2.1e}$$

Boundary conditions are:

at 
$$k = 0$$
;  $p_{rk} = p_r^0$ ;  $T_{rk} = T_r^0$ ;  $\rho_r = \rho_r^0$  (3.3.2.2a,b,c)

The pressure at the well datum used in evaluating equation 3.3.2.2a for production conditions is explicitly calculated at time plane n. Equations 3.3.2.1a-d are integrated over the length of the well riser,  $L_r$ , yielding the desired quantities  $p_r(L_r)$  and  $T_r(L_r)$ . The functions F and G are evaluated by the right-hand-side of equation 2.4.2.11 with the following equations used for calculating density and velocity:

$$\rho_{r} = \rho_{r}^{0} + \rho_{r}^{0} \beta_{p} (p_{r} - p_{r}^{0}) - \rho_{r}^{0} \beta_{T} (T_{r} - T_{r}^{0}) ; \qquad (3.3.2.3a)$$

$$\rho_{rk} v_{rk} = \rho_{r}^{0} v_{r}^{0} = \frac{\rho_{r}^{0} Q_{w}}{\pi r_{r}^{2}}.$$
 (3.3.2.3b)

The midpoint method of integration is a second-order method, which means that the error in  $p_r(L_r)$  and  $T_r(L_r)$  decreases as  $(\Delta \ell)^2$ . The extrapolation procedure improves the accuracy of the numerical integration by estimating results for  $p_r(L_r)$  and  $T_r(L_r)$  that would be obtained if the step length,  $\Delta \ell$ , were reduced to zero. Pressure and temperature at the end of the well riser are expressed by power-series expansions as a function of step length along the riser:

$$p_{r}(L_{r},\Delta \ell) = p_{r}(L_{r}) + \sum_{i=1}^{n} d_{pi} \Delta \ell^{2i}$$
 (3.3.2.4a)

$$T_r(L_r, \Delta \ell) = T(L_r) + \sum_{i=1}^{n} d_{Ti} \Delta \ell^{2i}$$
 (3.3.2.4b)

where

d are the coefficients in the series expansion for	
pressure (Pa/m); and	(3.3.2.4c)
d <sub>Ti</sub> are the coefficients in the series expansion for	
temperature (°C/m).	(3.3.2.4d)

Equations 3.3.2.4a and 3.3.2.4b can be written in vector form by defining:

$$\underline{\underline{Y}}(L_{r},\Delta \underline{x}) = \begin{bmatrix} p_{r}(L_{r},\Delta \underline{x}) \\ T_{r}(L_{r},\Delta \underline{x}) \end{bmatrix}$$
(3.3.2.5a)  
$$\underline{\underline{Y}}_{i} = \begin{bmatrix} d_{pi} \\ d_{Ti} \end{bmatrix}$$
(3.3.2.5b)

so that:

$$\underline{Y}(L_r, \Delta \ell) = \underline{Y}(L_r) + \sum_{i=1}^{m} \underline{Y}_i \Delta \ell^{2i}$$
(3.3.2.6)

The right-hand-side of equation 3.3.2.6 is approximated by a rational function,  $\underline{R}_{m}$ , that is, a quotient of two polynomials. The coefficients of the rational function are determined so that:

$$\underline{R}_{m}(L_{r},\Delta t_{j}) = \underline{Y}(L_{r},\Delta t_{j}) ; j = 0,1...m$$
 (3.3.2.7)

where

 $\Delta l_j$  is the spatial-step length for the jth integration from 0 to L (m).

Then, the desired solution,  $\underline{Y}(L)$ , is related to the approximating rational function by:

$$\underline{Y}(L_r) = R_m(L_r, 0)$$
 (3.3.2.8)

The algorithm is formed by defining  $R_m^j$  ( $\Delta \ell$ ) as the rational approximation which agrees with  $\underline{Y}(L_r, \Delta \ell)$  at  $\Delta \ell = \Delta \ell_j$ ,  $\Delta \ell_{j+1}$ , ...,  $\Delta \ell_{j+m}$ , where  $\Delta \ell_j > \Delta \ell_{j+1}$ , and defining  $R_m^j(0) = R_m^j$ . Then the  $R_m^j$  give better approximations to  $\underline{Y}(L_r)$  as j and (or) m increase. The extrapolation procedure is initiated by integrating equations (3.3.2.1 a-d) for a sequence of step lengths, L/2, L/4, L/6, L/8, ... to obtain values for  $R_0^{\circ}$ ,  $R_0^{-1}$ ,  $R_0^{-2}$ , .... Values of  $R_m^j$  for increasing j and m are calculated by the recurrence relation given in Gear, 1971, p. 95. The procedure is terminated when two successive approximations,  $R_k^{m-k}$  and  $R_k^{m-k+1}$ , are sufficiently close. The tolerance estimate for the fractional error can be set by the user with a default value of  $10^{-3}$ .

Depending on the rate of convergence, the step size may be increased or decreased for successive well-riser calculations. Sixth-order polynomials are the maximum order used for the rational approximation with a maximum of 10 different step sizes.

### 3.4 BOUNDARY CONDITIONS

All boundary conditions are specified on a cell-by-cell rather than on a zone-by-zone basis. The default-boundary condition is that of no dispersive or advective flux through the boundary faces of the cell. For a cell with three boundary faces, up to three different types of flux-boundary conditions can be applied, each to a different face. For example, a specified flux, an aquifer-influence function, and a leakage-boundary condition could be applied to the faces of a corner cell.

# 3.4.1 Specified Pressure, Temperature, and Solute-Mass Fraction

Specified-value boundary conditions are incorporated by replacing the flow and transport equations for those nodes, by equations of the form 'of equation 3.1.1c defining the specified values. These nodes could be removed from the set of simultaneous equations to be solved, by incorporating the known boundary values into the remaining equations; that has not been done in the present version of the HST3D simulator. For boundary conditions that change discontinuously with time, the value at time t<sup>n</sup> is taken to be the limit of the value at t<sup>n</sup> -  $\delta$ t, as  $\delta$ t + 0; that is, the jump in the boundarycondition value takes place after the time of change. This means that the effective value of a boundary condition over a time interval when a change occurs is the average value under centered-in-time differencing and the later value under backward-in-time differencing.

It should be noted that an initial hydrostatic-pressure boundary condition over depth will not be maintained under conditions of variable-density flow. Specification of hydrostatic-pressure boundary conditions over depth using a uniform initial density can cause disconcertingly large vertical flows to occur, when realistic fluid compressibility effects are incorporated during the simulation. Even if the compressibility is very small, the boundary pressure values need to be specified to four or five significant digits to "avoid vertical flows caused by roundoff error.

Since a specified-value boundary condition removes the equation for the corresponding variable (pressure, temperature, or mass fraction) from the set to be solved, some constraints do exist on what boundary conditions can be specified for a cell that has more than one boundary face. For example, if the pressure is specified, then the ability to specify a fluid flux, an aquifer-influence function, or a leakage boundary condition on the other boundary faces is lost.

#### 3.4.2 Specified-Flux Boundary Conditions

Discretization of the flow equations and transport equations causes the specified-flux boundary conditions to be incorporated as source terms in the finite-difference equations, as described by equation 3.1.1.25a and b. The specified fluxes are input as vector components at each of the respective boundary faces. Thus they are described on a cell-face basis, not by zone boundary. Fluid fluxes are input as volume fluxes; heat fluxes are input as energy fluxes; solute fluxes are input as mass fluxes.

Recall that a boundary cell can have up to three boundary faces, each with an outward normal vector pointing in one of the coordinate directions. The flux-vector components can specify flux only through a face whose normal is parallel to the vector component. Thus, the number of specified-flux vector components must be less than or equal to the number of boundary faces for a given cell. If the normal and the vector component point in opposite directions, flux is added to the boundary cell; if they point in the same direction, flux is withdrawn.

A persistent numerical error can arise in the case where only specifiedflux boundary conditions are employed for the entire region, because of the occurance of a zero eigenvalue for the discretized equation (Mitchell, 1969, p. 39-44). Errors generated by discontinuous changes in the boundary conditions with time or by discontinuities between the initial conditions and the boundary conditions will persist. If a specified-value boundary condition or 'flux-dependent-on-value boundary condition is applied over some part of the boundary, this problem vanishes, because the zero eigenvalue disappears.

A one-dimensional analysis shows that the integral form of derivation used for the specified-flux boundary conditions gives a discretization error of order  $\Delta t \Delta x$ . Thus, the finite-difference equations are only first-order accurate at the boundary cells in terms of specified flux.

#### 3.4.3 Leakage-Boundary Conditions

Leakage-boundary conditions are transformed into source-sink terms in a similar fashion to specified-flux conditions. They also are incorporated on a cell basis rather than on a zone basis. Equations 2.5.3.1.1a-c and 2.5.3.2.1a-c, when applied on a discrete grid, become for boundary cell, m:

$$Q_{Lm} = \frac{k_{Lm}}{\mu_{Lm}b_{Lm}} [(\rho_e \phi_e)_m - (p_m^n + \rho_m^n g z_m)$$
(3.4.3.1a)

- 
$$(\rho_e - \rho_m^u)g(z_e + z_m)/2] S_{BLm}$$

$$-\frac{k_{Lm}}{\mu_{Lm}b_{Lm}}S_{BLm}\delta p_{m};$$

 $Q_{Rm} = \gamma_R Q_{Lm};$ 

(3.4.3.1b)

where

 $Q_{Lm}$  is the volumetric flow rate at a leakage boundary (m<sup>3</sup>/s);  $Q_{Rm}$  is the volumetric flow rate at a river-leakage boundary (m<sup>3</sup>/s); and  $S_{BLm}$  is the part of the boundary cell surface that is a leakage boundary (m<sup>2</sup>).

The leakage-flow rate, of equation 3.4.3.1a, has an explicit term for the right-hand-side of the discretized system-flow equation, 3.1.4.1a, and an implicit factor for the left-hand-side.

3.4.4 Aquifer-Influence-Function Boundary Conditions

#### 3.4.4.1 Pot-Aquifer-Influence Function

The aquifer-influence-function boundary conditions for a pot aquifer are discretized by writing equation 2.5.4.1.1 for each cell face over which the pot-aquifer boundary condition applies. Let there be  $M_A$  pot-aquifer boundary condition cells. Then:

 $Q_{Am} = [\alpha_{be} + \varepsilon_e \beta_{pe}) \frac{\delta p_{Bm}}{\delta t} V_{em}, \quad m = 1, M_A; \quad (3.4.4.1.1)$ 

where

- <sup>2</sup>Bm is the rate of pressure change at the boundary of the inner region for cell m (Pa/s);
- vem is the volume of the outer-aquifer region that influences boundary cell m (m<sup>3</sup>); and
- Q<sub>Am</sub> is the volumetric flow rate across the boundary face for cell m between the inner- and outer-aquifer regions; (positive is into the inner region), (m<sup>3</sup>/s).

The volume of outer aquifer that influences boundary cell m usually is taken to be the permeability-weighted fractional area of the boundary face:

$$V_{em} = \frac{V_{e_{q=1}} \Sigma k_{mpq} S_{Ampq}}{M_{A} n_{p} 4};$$
  
$$\Sigma \Sigma \Sigma k_{mpq} S_{Ampq}$$
  
$$m=1 p=1 q=1$$

(3.4.4.1.2)

where

S<sub>Ampq</sub> is the area of the aquifer-influence function boundary face for cell m, face p, subdomain q (m<sup>2</sup>); and

 $k_{mpq}$  is the permeability for cell m, face p, subdomain q (m<sup>2</sup>).

This aquifer-influence-function flow rate gives only an implicit coefficient for the left-hand-side of the flow equation, 3.1.4.1a.

#### 3.4.4.2 Transient-Flow Aquifer-Influence Function

The transient-flow, aquifer-influence function is discretized by writing equations 2.5.4.2.6a-b for each cell at which this boundary condition applies. Thus, a different pressure history may occur at each boundary node. The aquifer-influence-function flow rate must be suitably apportioned among the boundary cell faces. The method used for HST3D is to make the fraction of the total flow rate that is apportioned to a given boundary cell the same as the ratio of that boundary-cell facial area to the total boundary facial area between the inner-and outer-aquifer regions. For cases where the inneraquifer region is strongly heterogeneous, apportionment by the product of hydraulic conductivity and facial area, using equation 3.4.4.1.2, would be more realistic. This would require modification to the program code.

The derivation of the flow rate given by equation 2.5.4.2.6a also was based upon a uniform pressure plus gravitational potential over the approximating cylindrical interface. A finite-difference flow simulation in the inner-aquifer region will yield a nonuniform distribution of pressures at the boundary nodes, except in special cases. In the numerical implementation of this aquifer-influence-function calculation, the pressure at each interfacial boundary node is taken to be the value computed by the discretized simulation calculation. This introduces an additional approximation, because any lateral or vertical flow in the outer-aquifer region, induced by the nonuniform pressure plus gravitational potential distribution over the interfacial boundary, is neglected.

Another approximation used is that the boundary between the inner- and outer-aquifer regions is represented by a cylindrical interface (fig. 2.5a); whereas, the actual boundary is a set of rectangular faces for the finitedifference discretization in cartesian coordinates of a three-dimensional, inner-aquifer region. In contrast, a two-dimensional, cylindrical gridding for the inner-aquifer region would have an exact cylindrical boundary. For a cartesian-coordinate system with the x-y axes horizontal, the equivalent radius,  $r_{I}$ , for the approximate interfacial boundary is calculated, so that the rectangular area and the equivalent circular area are the same; that is:

$$\pi r_{I}^{2} = (x_{Nx} - x_{1})(y_{Ny} - y_{1}) . \qquad (3.4.4.2.1)$$

Equation 3.4.4.2.1 will be a poor approximation for long, slender rectangular areas in the x-y plane. For boundaries between the inner- and outer-aquifer regions that do not completely surround the inner aquifer laterally, the apportionment factor  $\gamma_{Am}$ , must contain an angle-of-influence factor,  $f_A$ . Then:

$$\gamma_{Am} = f_{\theta} S_{Ampq} ; \qquad (3.4.4.2.2)$$

where

SAmpq is the area of the aquifer-influence function boundary face for cell m, face p, subdomain q (m<sup>2</sup>); and f<sub>θ</sub> is the angle-of-influence factor for the simulation region (-). This factor is the fraction of a full circle that the boundary between the inner- and outer-aquifer regions subtends. For example, an outer-aquifer region that surrounds half of the inner-aquifer simulation region (fig. 2.5b) would have an angle-of-influence factor of  $\frac{1}{2}$ .

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The Carter-Tracy approximation is used to avoid successive recomputation of the convolution integral that gives the flow rate at the transient aquiferinfluence-function boundary. After discretization of time, the expression for the flow rate across the boundary between the inner-aquifer region and the outer-aquifer region is [Kipp (1986)]:

$$Q_{Am}^{n+1} = Y_{Am} \frac{2\pi k_{e} b_{e}}{\mu_{e}} \left[ \frac{P_{m}^{n} - P_{m}^{0} - \frac{dP_{U}}{dt}}{P_{U}^{n+1} - \frac{dP_{U}}{dt}} \right|^{n+1} \frac{V_{m}^{n}}{2\pi r_{I}^{2} (\alpha_{be} + \varepsilon_{e} \beta_{pe}) b_{e}} + \delta p_{m}}{P_{U}^{n+1} - \frac{dP_{U}}{dt}} \right]^{n+1} t^{n}$$
(3.4.4.2.3)

The Carter-Tracy approximation is based on representing the continuous flow rate of equation 2.5.4.2.6a by a discontinuous sequence of constant flow rates so that the cumulative net inflow from the start of the simulation to the given time is the same for the convolution integral and the current constant-flow rate. This approximation is exact for constant-flow rates. Therefore, slowly varying flow rates are more accurately handled by the Carter-Tracy approximation than rapidly varying ones. The major disadvantage of the Carter-Tracy approximation is the inaccuracy of the computation of the discretized flow rate in the case where boundary-flow rates vary with time. Effects on the computed-flow rate appear as a significant time lag and smoothing of transients. Errors can be serious for boundary-flow rates whose variations are large relative to the average value.

Note that the computer storage requirements for this calculation are only the cumulative flow,  $W_m^n$ , the pressure at the end of the nth time step,  $p_m^n$ , and

the flow-rate allocation factor,  $\gamma_{Am}$ , for each node on the boundary between the inner- and outer-aquifer regions. The values of  $p_m^n$  are from the flow simulation, so the additional storage amounts to only two times the number of AIF boundary nodes.

Equation 3.4.4.2.3 is of the form:  

$$Q_{Am}^{n+1} = a_1(t') + a_2(t') \delta p_m;$$
 (3.4.4.2.4)

where  $a_1$  is the known flow rate term added to the right-hand side of the discretized flow equation for node m; and  $a_2$  is the implicit term added to the left-hand side factor.

The values of  $P_U'(t')$  and  $\frac{dP_U'}{dt'}$  are obtained usually from tables by inter-

polation. Values of the dimensionless pressure function in response to a unit-withdrawal flow rate at the AIF boundary have been tabulated by Van Everdingen and Hurst (1949) for the infinite cylindrical region and for regions with a finite outer-boundary radius.

However, it is much more convenient to use the approximate analytical representation developed by Fanchi (1985). He employed linear regression analysis to obtain the following equation that approximates the Van Everdingen and Hurst (1949) aquifer influence functions with very small errors. In the notation of this report;

$$P_{U}'(t') = b_0 + b_1 t' + b_2 \ln(t') + b_3 \ln^2(t') \qquad (3.4.4.2.5)$$

Table 3.2 adapted from Fanchi (1985) contains values of the  $b_i$  coefficients for several cases. The first line gives the coefficients for the analytical approximation to equation 2.5.4.2.5b for the case of an infinite outer-aquifer region. The subsequent lines are for various values of R for the case of a finite outer-aquifer region with no flow at the exterior boundary, where R is the ratio of exterior to interior radius for the outer-aquifer region. As with the previous boundary conditions, the heat- and solute-advective transport rates across the AIF boundary are calculated by multiplying the flow rate by the appropriate density, enthalpy, and mass-fraction values, depending on the direction of flow.

R	PO	<i>b</i> 1	<sup>b</sup> 2	<sup>b</sup> 3
œ	-0.82092	3.68×10-4	-0.28908	-0.02882
.5	-0.10371	-1.66657	0.04579	0.01023
.0	-0.30210	-0.68178	0.01599	0.01356
.0	-0.51243	-0.29317	-0.01534	0.06732
0	-0.63656	-0.16101	-0.15812	0.09104
0	-0.65106	-0.10414	-0.30953	0.11258
.0	-0.63367	-0.06940	-0.41750	0.11137
.0	-0.40132	-0.04104	-0.69592	0.14350
.0	-0.14386	-0.02649	-0.89646	0.15502

Table 3.2--Coefficients for the analytical approximations to the Van Everdingen and Hurst aquifer-influence functions

#### 3.4.5 Heat-Conduction Boundary Condition

The heat-conduction boundary condition is a flux-type boundary condition, with the heat flux dependent on the thermal parameters and the thermal history at the boundary, and in the conducting medium, which lies outside the simulation region. Equation 2.5.5.7 gives the heat flux that is applied on a cell basis to the source term in the heat-transport equation. Finite-difference approximations to equations 2.5.5.3a-c and 2.5.5.4a-c are solved at each time step for each heat-conduction, boundary-condition cell. Central-differencing in time is used giving a tridiagonal-matrix equation to be solved numerically. The user specifies the spatial mesh extending out from the boundary into the conducting medium. Up to 10 nodes are allowed with variable spacing. The first node must be on the boundary. The Thomas algorithm (Varga, 1962, p. 195) is used to obtain the numerical solutions for  $T_e$  and  $T_U$ . Then the heat flux for cell m is given by:

$$Q_{HCm} = -K_{em} \left[ \frac{T_{e}(z_{n2}) - T_{e}(z_{n1})}{z_{n2} - z_{n1}} + \frac{T_{U}(z_{n2}) - T_{U}(z_{n1})}{z_{n2} - z_{n1}} \right] \delta T_{Bm} S_{BHCm}; (3.4.5.1)$$

where

z<sub>n1</sub> and z<sub>n2</sub> are the first two nodes moving in the outward normal direction from boundary cell m (m).

Node z is coincident with the boundary node of the simulation region, so the boundary temperature is:

$$T_{Bm} = T_{m}$$
 (3.4.5.2)

Equation 3.4.5.1 is of the form:

.

 $Q_{HCm} = a_1(t) + a_{2m}(t) \delta T;$  (3.4.5.3)

where the  $a_1(t)$  term goes into the source term for the cell heat-transport equation, and the  $a_2(t)$  term goes into the thermal-coefficient matrix for cell m. An initial temperature profile can be specified for the heat-conduction region. However, the same profile is used for all cells with a heat-conduction boundary condition.

3.4.6 Unconfined-Aquifer, Free-Surface Boundary Condition

The unconfined-aquifer, free-surface boundary condition is implemented by modifying the pressure-coefficient terms in the discretized equations for flow and solute-transport, and adjusting the fluid volume or saturated thickness of the uppermost layer of cells in the simulation region. A free surface is not allowed when the heat-transport equation is being solved, because no satisfactory method has been developed to handle the conductive-heat flux through the free-surface boundary that moves with time through the unsaturated porous medium.

The location of the free surface within an upper-boundary cell is established by linearly extrapolating the nodal pressure to the elevation of zero (atmospheric) pressure, using the fluid density in that cell. The fraction of the cell thickness that is saturated is given by:

$$f_{FS} = 1 + \frac{p_m}{\rho_m g_2^1 (z_{NZ}^2 - z_{NZ-1}^2)};$$
 (3.4.6.1)

where

 $f_{FS}$  is the fraction of the cell thickness that is saturated (-); and  $p_m$  is the pressure at node m (Pa);

 $z_{N_{\mathcal{T}}}$  is the elevation of the upper boundary (m); and

z<sub>Nz-1</sub> is the elevation of the next layer of nodes down from the upper boundary (m).

This equation is valid for coordinate systems with the z-axis pointing vertically upward. Remember that the pressure value is relative to atmospheric pressure, so that atmospheric pressure does not appear explicitly in equation 3.4.6.1. This fraction is allowed to range from zero to two; that is, the free surface is allowed to rise above the upper boundary of the simulation region to a distance that is equal to the upper-layer half-cell thickness. This rise is effectively the same as using full cells in the vertical direction for the upper layer when a free-surface boundary condition is being used. The extra cell height allows for a greater variation of the free surface than is possible with the normal cell height in the upper layer.

When designing the grid for a free-surface boundary problem, the uppermost layer of cells must be made thick enough to accommodate the maximum variations in the free-surface location. With the present algorithm, the free surface may not drop below the lower boundary of the uppermost layer of cells. No conversion to confined flow conditions is made if it does rise above the extended upper boundary. These restrictions can be burdensome if a large drawdown cone is created by a well pumping an unconfined aquifer, because the uppermost layer of cells may need to be so thick that vertical gradients are represented poorly. The present version of the HST3D program is suitable for simulation of only modest drawdowns relative to aquifer thickness for unconfined conditions.

If desired by the user, a message may be printed when the free surface rises above the extended upper boundary of the region or falls below the bottom of the uppermost layer of cells, or cells in a lower layer become partially saturated.

To obtain the appropriate coefficients for the flow equation (3.1.4.1a) at the free-surface boundary cells, we evaluate the terms of equation 3.1.1.8, this time, including the saturation fraction of equation 3.4.6.1, using equation 3.1.3.5, and assuming that the porosity is constant, the fluid compressibility is zero and isothermal conditions exist, to obtain:

$$C_{33} = \left[\sum_{s=1}^{8} \sum_{s=1}^{N} \sqrt{\frac{1}{2}g(z_{Nz} - z_{Nz-1})\delta t}\right]; \qquad (3.4.6.2a)$$

and the coefficient  $C_{31}$  remains unchanged from equation 3.1.4.1d.

Using the same procedure as for equations 3.4.6.2a and 3.4.6.2b, the corresponding terms for the solute-transport equation (3.1.4.4a) are:

$$C_{11} = \sum_{s=1}^{8} (\varepsilon_{s} + \rho_{b}K_{d})_{s} V_{s}[\rho_{m}^{n+1} f_{Fs}^{n+1} + w_{m}^{n} \rho_{o}\beta_{w}]/\delta t; \quad (3.4.6.3a)$$

 $C_{13} = \left[\sum_{s=1}^{8} (\varepsilon_{s} + \rho_{b}K_{d})_{s} V_{s} - \frac{V_{m}^{n}}{\frac{1}{2g(z_{Nz} - z_{Nz-1})}}\right] / \delta t . \qquad (3.4.6.3b)$ 

Additional terms arise from the source-sink term in the solute equation that are functions of solute-mass fraction. They form part of the  $C_{11}$  and  $C_{13}$ coefficients. It has also been assumed that all of the solute in a cell is either in the fluid phase or is sorbed on the saturated part of the porous medium. No account is taken of solute that might sorb onto the porous medium and be left behind, when the free surface falls. This simplification is consistent with this approximate treatment of a free-surface boundary condition. The other terms in the flow and transport equations have the saturated fraction parameter included, as necessary, for the cell facial-area terms involving the x and y directions. No additional contributions to the  $C_{ij}$  terms occur, because the dispersive and advective coefficients are evaluated at time t<sup>n</sup> only.

The case of a free-surface boundary with accretion of fluid by infiltration is also handled in an approximate fashion. The fluid flux is specified at the upper boundary of the cell, and the associated temperature and mass fraction determine the amount of heat and solute that enter through the free surface.

3.5 INITIAL CONDITIONS

and

The numerical implementation of the initial conditions is straightforward. Values of pressure, temperature and mass fraction are set to the initial value distributions for each node in the simulation region given in the form of equation 3.1.1b, that is:

at t=0, 
$$u_{ijk} = u_{ijk}^0$$
.

(3.5.1)

The specified distributions can vary on a node-by-node basis or be zones of constant conditions. Available options for the initial pressure distribution include hydrostatic equilibrium, a water-table surface, or a pressure field specified node by node. The water-table surface is specified for the upper layer of cells only. These initial distributions are based on the initial temperature and solute-mass fraction distributions.

The hydrostatic-equilibrium pressure distribution takes the fluid compressibility into account. The calculation proceeds from the bottom of the region upward or from the top of the region downward depending on the elevation of the specified initial pressure.

The water-table elevation surface is specified for the upper layer of nodes only. Hydrostatic equilibrium is assumed to compute the pressure distribution elsewhere in the simulation region. When specifying the initial pressure field on a node-by-node basis, it is permissible to include nodes that are outside the simulation region. This is for ease of data input by rectangular zones or by ascending node number.

## 3.6 EQUATION SOLUTION

Equations 3.1.4.1a, 3.1.4.2a and 3.1.4.4a can be written in matrix form as:

$$\begin{bmatrix} C_{11} & C_{12} & C_{13} \\ C_{21} & C_{22} & C_{23} \\ C_{31} & C_{32} & C_{33} \end{bmatrix} \begin{bmatrix} \delta w_{m} \\ \delta T_{m} \\ \delta p_{m} \end{bmatrix} = \begin{bmatrix} E_{11} & 0 & E_{13} \\ 0 & E_{23} & E_{23} \\ 0 & 0 & E_{33} \end{bmatrix} \begin{bmatrix} \delta w_{m} \\ \delta T_{m} \\ \delta p_{m} \end{bmatrix} + \begin{bmatrix} F_{1} \\ F_{2} \\ F_{3} \end{bmatrix}$$
(3.6.1a)  
$$\begin{bmatrix} C_{11} & C_{12} & C_{13} \\ \delta T_{m} \\ \delta p_{m} \end{bmatrix} + \begin{bmatrix} F_{1} \\ F_{2} \\ F_{3} \end{bmatrix}$$
(3.6.1b)  
$$= \begin{bmatrix} R_{1} \\ R_{2} \\ R_{3} \end{bmatrix} ;$$

where

 $\frac{E_{ij}}{\delta p_m}$  are the coefficient vectors in the discretized equations;  $\frac{\delta p_m}{\delta p_m}$  is the change in pressure for node m (Pa);  $\frac{\delta p_m}{\delta p_m}$  is the change-in-pressure vector for node m (Pa);  $\frac{\delta T_m}{\delta T_m}$  is the change in temperature for node m (°C);  $\frac{\delta T_m}{\delta m}$  is the change-in-temperature vector for node m (°C);  $\frac{\delta w_m}{\delta m}$  is the change in mass fraction for node m (-);  $\frac{\delta w_m}{\delta m}$  is the change-in-mass-fraction vector for node m (-); and

 $F_i$  are the known terms at time n in the discretized system equations.

The vectors of the changes in the dependent variables contain the values for each node connected to node m plus node m itself. The <u>E</u> vectors in equation 3.6.1a have seven components each that correspond to node m and its six neighbors in the three coordinate directions. The known terms  $F_i$  are those that do not contain  $\delta w$  for i=1,  $\delta T$  for i=2, or  $\delta p$  for i=3. The terms  $E_i$  and  $F_i$  can be functions of pressure, temperature, and mass fraction at time n which gives explicit linking of the three equations. Implicit linking is through the <u>C</u> matrix on the left-hand side. The equations are written in reverse order to what has been done previously, with 1 referring to the solute-transport equation, 2 referring to the heat-transport equation, and 3 referring to the flow equation. Equation 3.6.1b is written for each node in the simulation region, giving a set of 3M simultaneous equations to be solved for the unknown vectors,  $\delta p$ ,  $\delta T$  and  $\delta w$ , where M is the total number of nodes in the region.

3.6.1 Modification of the Flow and Transport Equations

To avoid storing a  $3M \times 3M$  matrix and vectors of length 3M, a sequential solution scheme has been developed by Coats and others (1974) and was used by INTERCOMP Resource and Development and Engineering, Inc. (1976). This algorithm consists of solving a modified flow equation then a modified heat-transport equation, then the solute-transport equation in turn for each time step. The modified equations are obtained by a partial Gauss reduction

of equation 3.6.1b transforming the capacitance matrix,  $\underline{C}$ , into uppertriangular form. Thus we have:

 $\begin{bmatrix} C_{11} & C_{12} & C_{13} \\ 0 & C_{22} & C_{23} \\ 0 & 0 & C_{33} \end{bmatrix} \begin{bmatrix} \delta w_{m} \\ \delta T_{m} \\ \delta p_{m} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ C_{24} & 1 & 0 \\ C_{34} & C_{35} & 1 \end{bmatrix} \begin{bmatrix} R_{1} \\ R_{2} \\ R_{3} \end{bmatrix} ;$   $C_{22}' = C_{22} - \frac{C_{12}C_{21}}{C_{11}} ;$  (3.6.1.1b)

where

:

$$C_{23}' = C_{23} - \frac{C_{13}C_{21}}{C_{11}};$$
 (3.6.1.1c)

$$C_{24} = -\frac{C_{21}}{C_{11}};$$
 (3.6.1.1d)

$$C_{33} = C_{33} - \frac{C_{13}C_{31}}{C_{11}} - \frac{(C_{23}C_{11} - C_{13}C_{21})(C_{32}C_{11} - C_{12}C_{31})}{C_{11}(C_{22}C_{11} - C_{12}C_{21})}; (3.6.1.1e)$$

$$C_{34} = -\frac{C_{31}}{C_{11}} + \frac{C_{21}(C_{32}C_{11} - C_{12}C_{31})}{C_{11}(C_{22}C_{11} - C_{12}C_{21})}; \qquad (3.6.1.1f)$$

$$C_{33} = - \frac{C_{32}C_{11} - C_{12}C_{31}}{C_{22}C_{11} - C_{12}C_{21}}$$
(3.6.1.1g)

### 3.6.2 Sequential Solution

The finite-difference approximations (equation 3.6.1.1a) to the groundwater flow, heat transport, and solute transport equations are solved sequentially. First, the flow equations are solved for the pressures. Then the

pressures are used to update the coefficients in the heat equations and back substitute for the op terms on the left-hand side. Second, the heat equations are solved for the temperatures. The temperatures and pressures are used in the solute equation to update the coefficients and to back substitute for the op and oT terms on the left-hand side. Finally the solute equations are solved for the mass fractions, which completes an iteration. Thus only M equations at one time are being solved with a coefficient matrix that is  $M \times M$ in size. M is the total number of active nodes in the simulation region. Actually the storage requirement is reduced by taking advantage of the matrix sparsity as will be described later. The solution cycle is repeated until convergence is achieved, that is, when the fractional change in fluid density in each cell is less than a tolerance value. The default value is 0.001 fractional change in density. Since changes in both temperature and mass fraction cause changes in density, a secondary tolerance criterion is set to determine whether the heat equation or the solute equation or both equations must be solved again. The secondary tolerance is 0.0005 fractional change in density. If the maximum density change due to temperature changes after solution of the heat equation or due to mass-fraction changes after solution of the solute equation is less than the secondary tolerance, then that equation is excluded from the iterative cycle for the remainder of the calculations for that time step. However, a final solution of the excluded equation is performed after convergence of the iterative cycle for the two remaining equations. In practice, convergence is usually achieved within three iterations for a given time step. Lack of convergence may indicate that the time step is too large.

Equation 3.6.1.1a is transformed into:

 $\Delta \cdot \underline{\delta u} = \mathbf{b};$ 

(3.6.2.1a)

where

 $\frac{\delta u}{\delta T} = \begin{bmatrix} \frac{\delta w}{\delta T} \\ \frac{\delta T}{\delta p} \end{bmatrix};$ 

(3.6.2.1b)

by shifting all the terms on the right-hand side that contain op, oT or ow to the left-hand side. Equation 3.6.2.1a is a linear-matrix equation for the region that can be solved using techniques to be described in section 3.7.

### 3.7 MATRIX SOLVERS

The linearized flow and transport finite-difference equations are solved in turn by one of the solution algorithms for linear, sparse-matrix equations. For the present version of the simulator, two such algorithms are available. One is a direct equation solver that uses Gauss elimination, after reordering the equations for a savings in computation time and computer-storage requirements. Alternating diagonal planes are used for the reordering. This method is referred to as the D4 solution technique; it was developed by Price and Coats (1974).

The other sparse-matrix equation solver uses the two-line, successiveoverrelaxation method. This is one of a class of block-iterative methods described by Varga (1962, p. 199-200). In this solver, two lines of nodes along a selected coordinate direction are solved together by direct elimination. One iteration sweep consists of solving for the nodal values for each pair of lines, plus the odd left-over line, if necessary. Overrelaxation is used to speed convergence, and the optimum overrelaxation factor is estimated, using the eigenvalue estimation technique of Varga (1962, p. 2£4-288) at the beginning, and then, every n time steps, as specified by the user. In the process of estimating the optimum-overrelaxation factor, the solution is tested in all three coordinate directions, and the direction with the bestconditioned iterative matrix is selected. It may be different for each of the three equations.

The form of the equations to be solved is the same as equation 3.6.2.1a, but now the matrix  $\underline{A}$  is a sub-matrix of the original one, containing the coefficients of only the flow, or heat transport, or solute transport equations. Matrix  $\underline{A}$  has the banded structure under the original nodal numbering scheme, where the index i is incremented first; the index j is

incremented second; and the index k is incremented third. Figure 3.7 shows the structure of the matrix  $\underline{A}$ , with the bandwidths given by N<sub>X</sub> N<sub>y</sub> and N<sub>z</sub> where  $\boldsymbol{N}_i$  is the number of nodes in the ith direction. The rectangular prism of nodes also is shown in figure 3.7, which encompases the entire simulation region.

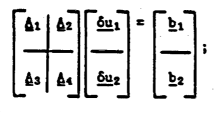
## 3.7.1 The Alternating Diagonal, D4, Direct-Equation Solver

The alternating diagonal reordering scheme was developed by Price and Coats (1974). In three dimensions, the equations are grouped by diagonal planes of nodes. A plane is defined by a fixed sum of the nodal subscripts. designated by the index, m, so:

$$i + j + k = m; m = 3, 4..., H;$$
 (3.7.1.1a)

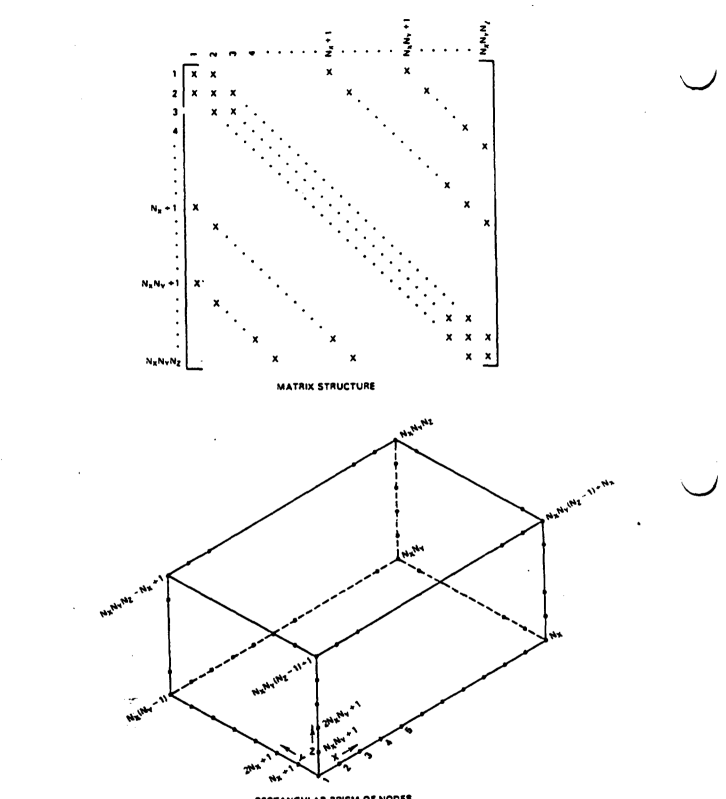
where  $M = N_x + N_y + N_z$ .

If M is even, then the order of plane-index selection for reordering of the node numbers should be m = 3,5,7,...M-1,4,6,8...M. If M is odd, then the order should be m = 3,5,7,...M,4,6,8,...M-1. For each plane index, m, the points should be numbered in order of decreasing k, decreasing j, and increasing i, assuming  $N_x > N_y > N_z$ . Any excluded cells are skipped during the node renumbering. The matrix A under D4 ordering takes the form shown in figure 3.8. This matrix can be partitioned as shown, so



(3.7.1.2)

(3.7.1.1b)



RECTANGULAR PRISM OF NODES

Figure 3.7.--Sketch of the matrix structure and the rectangular prism of nodes of the flow or heat-transport, or solute-transport equation in finite-difference form.

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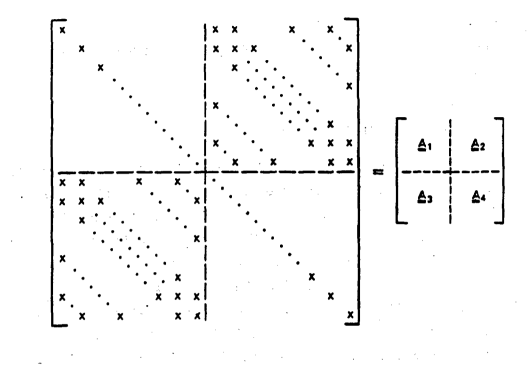


Figure 3.8.--Sketch of the matrix structure with the D4 alternatingdiagonal-plane, node-renumbering scheme.

where

•

 $\underline{A}_1$  and  $\underline{A}_4$  are diagonal and  $\underline{A}_2$  and  $\underline{A}_3$  are sparse matrices.

1.

Forward elimination gives:

$$\begin{bmatrix} \underline{\underline{A}_1} & \underline{\underline{A}_2} \\ \underline{\underline{Q}} & \underline{\underline{A}'_4} \end{bmatrix} \begin{bmatrix} \underline{\underline{\delta}\underline{u}_1} \\ \underline{\underline{\delta}\underline{u}_2} \end{bmatrix} = \begin{bmatrix} \underline{\underline{b}_1} \\ \underline{\underline{b}'_2} \end{bmatrix};$$

(3.7.1.3)

where

 $\underline{A}_4$  is a band matrix with a maximum bandwidth which is the same as for the original matrix  $\underline{A}_1$ .

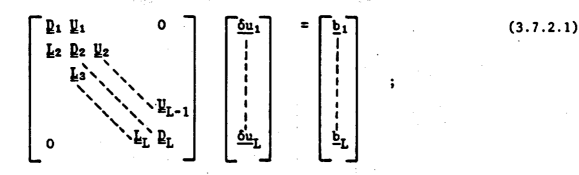
The solution for  $\underline{\delta u_2}$  is obtained by standard Gaussian elimination. Back substitution is used to compute  $\underline{\delta u_1}$  by:

$$\underline{\delta u_1} = \underline{A_1}^{-1} \underline{b_1} - \underline{A_1}^{-1} \underline{A_2} \underline{\delta u_2}. \tag{3.7.1.4}$$

The work required with the D4 reordering is from 17 to 50 percent, and the storage is from 33 to 50 percent of that using standard ordering. When the D4 ordering is selected with direct Gaussian elimination, the renumbering is entirely transparent to the user. The storage requirement for the  $\underline{A}_4$  matrix is minimized by employing variable bandwidth storage (Jennings, 1977, p. 97). The matrix is stored by rows, with the length of each row being sufficient to accommodate the fill-in that occurs during elimination. Two pointer arrays are necessary: one that contains the indices of the diagonal elements, and the other that gives the bandwidth to the right of the diagonal. The next row begins in the location just after that specified by the diagonal-element location, plus the right-side bandwidth.

### 3.7.2 The Two-Line, Successive-Overrelaxation Technique

This iterative matrix-equation solution technique, abbreviated L2SOR, is a block successive-overrelaxation algorithm as described by Varga (1962, sec. 6.4) or Jennings (1977, p. 202). Equation 3.6.2.1a for only the flow or heat transport or solute transport equation may be written with a partitioned <u>A</u> matrix (fig. 3.9) as:



where the sparse submatrices  $\underline{P}_k$  are penta-diagonal and the submatrices  $\underline{L}_k$  and  $\underline{U}_k$  are diagonal (fig. 3.9). There are L pairs of mesh lines with the last set containing only the odd remaining line, if one exists. These matrices are of size  $2N_x \times 2N_x$ , where the nodes have been numbered in the normal way, and the two lines have been selected to be in the x-direction. Some rearrangement would be necessary, if the y or z-directions were selected, but the basic structure would be the same. Then the iterative technique is expressed by the following equations:

$$[\underline{p}_{\ell} + \underline{w}_{\ell}] \underline{\delta u}_{\ell} \overset{\vee+1}{=} [-\underline{w}_{\ell} \underbrace{U}_{\ell} + (1-\underline{w}) \underline{p}_{\ell}] \underline{\delta u}_{\ell} \overset{\vee}{+} \underline{w}_{\ell}; \ell = 1, 2...L; (3.7.2.2)$$

where

v is the iteration counter; and

w is the overrelaxation factor.

Direct elimination is used to solve equation 3.7.2.2, after a renumbering is performed transverse to the direction of the two-lines. This renumbering compresses the bandwidth so no fill-in occurs. The iterations could be terminated when a vector-difference norm is less than a specified tolerance (Jennings, 1977, p. 184). That is, when:

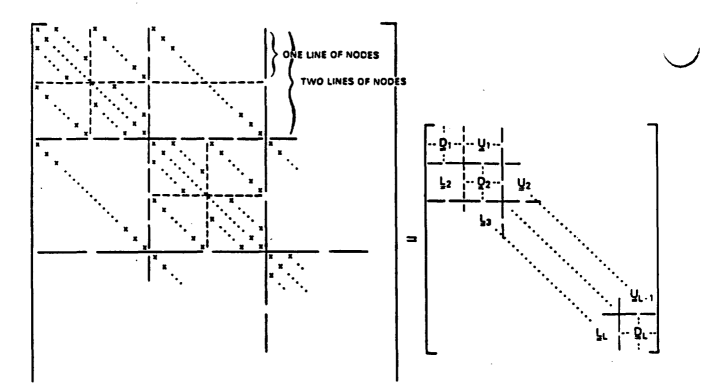


Figure 3.9--Sketch of the matrix structure with the two-line, successiveoverrelaxation, node-renumbering scheme.

$$\max_{m} \frac{\left| \delta u_{m}^{\nu+1} - \delta u_{m}^{\nu} \right|}{\left| \delta u_{m}^{\nu+1} \right|} \leq \varepsilon_{SOR}; m = 1, 2...M; \qquad (3.7.2.3)$$

where

\$\sum\_{SOR}\$ is the specified tolerance; and
m is the node number.

An alternate termination criterion is from Remson and others (1971, p. 185). Properties of the iteration matrix are taken into account by terminating the iteration when:

$$\frac{R(L_{\omega})}{1-R(L_{\omega})} \qquad \frac{\left| \begin{array}{c} \delta u_{m}^{\nu+1} - \delta u_{m}^{\nu} \right|}{\left| \begin{array}{c} \delta u_{m}^{\nu+1} \right|} \right| \leq \varepsilon_{SOR}; m = 1, 2...H; \qquad (3.7.2.4a)$$

where

 $R(L_{ij})$  is the spectral radius of the SOR iteration matrix.

Now from Varga (1962, p. 111):

1 1

$$R(L_{\rm w}) = w_{\rm opt} - 1,$$

where

w is the optimum overrelaxation factor.

This alternate criterion is used in the HST3D simulator. The tolerance,  $\varepsilon_{\rm SOR}$ , is set to 1 × 10<sup>-5</sup> by default, but it may be changed by the user.

The optimum overrelaxation factor is determined from estimates on the eigenvalues of the associated block Gauss-Seidel matrix. This is a combination of the power method and Perron-Frobenius theory of nonnegative matrices, presented by Varga (1962, p. 283-288). The algorithm is as follows: A two-line, Gauss-Seidel solution of the matrix equation:

$$\underline{A} \, \underline{\delta u} = \underline{0} ; \qquad (3.7.2.5a)$$

is performed with a starting vector of unit components, that is,

$$\underline{\delta u}_{\underline{\ell}}^{\nu+1} = [\underline{p}_{\underline{\ell}} + \underline{L}_{\underline{\ell}}]^{-1} [-\underline{U}_{\underline{\ell}}] \underline{\delta u}_{\underline{\ell}}^{\nu} ; \underline{\ell} = 1, 2... L \qquad (3.7.2.5b)$$

with 
$$\delta u_0^o = [1]$$
. (3.7.2.5c)

Then the minimum and maximum estimates of the dominant eigenvalue for  $\underline{A}$  are:

$$\lambda_{\min}^{\nu} = \frac{\min}{i} \frac{\delta u_i^{\nu+1}}{\delta u_i^{\nu}}; \qquad (3.7.2.6a)$$

(3.7.2.4b)

and 
$$\lambda_{\max}^{\nu} = \max_{i}^{\max} \frac{\delta u_{i}^{\nu+1}}{\delta u_{i}^{\nu}};$$
 (3.7.2.6b)

which satisfy

$$\lambda_{\min}^{\upsilon} < \lambda_{\min}^{\upsilon+1} < R(L_1) < \lambda_{\max}^{\upsilon+1} < \lambda_{\max}^{\upsilon}; \qquad (3.7.2.6c)$$

1

where

 $R(L_1)$  is the spectral radius of the Gauss-Seidel iteration matrix,  $[\underline{D}+\underline{L}]^{-1}[-\underline{U}]$ . Now, since:

$$w_{opt} = \frac{2}{1 + (1 - R(L_1))^2};$$
 (3.7.2.7)

let

$$\omega_{\min}^{\nu} = \frac{2}{1 + (1 - \lambda_{\min}^{\nu})^{\frac{1}{2}}}; \qquad (3.7.2.8a)$$

and

$$w_{\max}^{v} = \frac{2}{1 + (1 - \lambda_{\max}^{v})^{\frac{1}{2}}}$$
 (3.7.2.8b)

Then it can be shown that for the matrices,  $\underline{A}$ , that arise from the finite-difference equations, which are diagonally dominant and irreducible (Varga, 1962, p. 177):

$$w < w^{\nu+1} < w < w^{\nu+1} < w$$
  
min min opt max max;

/ where

 $w_{ont}$  is the optimum overrelaxation factor.

For  $\underline{A}$  nonsymmetric, the mesh spacing must be sufficiently small to ensure diagonal dominance. One requirement for irreducibility is that equations for nodes with specified value boundary conditions are eliminated from the set that forms  $\underline{A}$ . This elimination is not done in the present version of HST3D, and this sometimes causes difficulties in calculating  $w_{opt}$ , because a reducible matrix can have a maximum eigenvalue that is zero. The eigenvalues of the Gauss-Seidel iteration matrix must be real and nonnegative for the optimum-overrelaxation-factor calculation formula, equation 3.7.2.7, to apply.

The iterations are terminated when:

$$\frac{\omega_{\max} - \omega_{\min}}{2 - \omega_{\max}} \leq 0.2 .$$
 (3.7.2.10)

This algorithm provides an estimate of the optimum overrelaxation factor with rigorous upper and lower bounds before the iterative solution scheme is begun. Since the matrix  $\underline{A}$  changes with time, recomputation of the estimate of  $w_{opt}$  is performed every n time steps, as set by the user (default is 5). The power method algorithm converges when the dominant eigenvalue of  $\underline{A}$  is real, and it converges more quickly the smaller the ratio of the second dominant eigenvalue to the dominant eigenvalue. The convergence rate to  $w_{opt}$  can be discouragingly slow when this ratio is near one.

(3.7.2.9)

#### 3.7.3 Choosing the Equation Solver

The choice of equation solver depends on the size of the problem and the rate of convergence of the iterative method. The minimum half bandwidth of matrix  $\underline{A}$  is the product of the two smaller numbers of nodes,  $N_x$ ,  $N_y$ , and  $N_z$ . For a half bandwidth of 50, the work of the direct D4 solver is equivalent to <sup>c</sup> about 68 iterations of the L2SOR solver (Price and Coats, 1974). The other <sup>c</sup> consideration is the greater storage requirement of the D4 method, because of the pointer arrays required and the partial fill-in of the  $\underline{A}_4$  matrix that occurs. The L2SOR method causes no extra fill-in of nonzero elements in the  $\underline{A}$ matrix; however, rapid convergence rates are highly dependent on calculation of an accurate optimum overrelaxation factor. It is difficult to give any general rule for selection of matrix solver. Storage requirements probably will determine the best choice. The HST3D program writes out the storage requirements for both methods at the beginning of the simulation.

### 3.8 GLOBAL-BALANCE CALCULATIONS

Global balances for fluid mass, enthalpy, and solute mass are calculated at the end of each time step. Cumulative totals as well as increments over "each time step are computed. The primary use of the global-balance calculations is to aid in the interpretation of the magnitudes of transport that are occurring in the simulated system, and their distribution among the various types of boundary conditions and sources. Each of the system equations represents a mass or energy balance over each cell. By summing over the cells, we obtain the global-balance equations that relate the total change of mass or energy to the net boundary flow rates and the net injection through wells. The solute balance includes sorption and disappearance by reaction. The global-balance equations are integrated over the current time step to obtain the incremental changes.

The fluid-flow, heat-flow, and solute-flow rates at specified-value boundary nodes are obtained by evaluating the appropriate system equation at each specified-boundary-value cell and computing the flow of fluid, heat, or

solute across the region boundary for that cell, necessary to satisfy the fluid-balance, heat-balance, or solute-balance equation. Heat and solute fluxes that result from cross-derivative terms are neglected for the heatbalance and solute-balance equations for specified temperature and specified mass-fraction cells. The temporal differencing method is taken into account, also. This means that, for example, if the specified boundary pressure changes over a time step, and centered-in-time differencing is used, then the pressure at the beginning of the time step is effective over the first half of that time step, and the pressure at the end of the time step is effective over the second half. Therefore, the flow rate induced by the new specified pressure is effective for only half the time step. The appropriate equation for the boundary-flow rate calculation is the flow equation for specifiedpressure boundary cells, the heat-transport equation for specified-temperature boundary cells, and the solute-transport equation for specified-mass fraction boundary cells. Thus, the cell-balance equations are satisfied exactly for specified-value boundary cells.

The fluid, heat, or solute residual is defined as the change in the amount of the quantity present, minus the net flow of that quantity into the region. Thus, a positive residual means that there is an excess of that quantity present over what would be expected based on the net flows over the time interval. The various flows, amounts present, and residuals are printed in tabular form. Fractional residuals are defined as a ratio of the residual to the larger of the inflow, outflow, or accumulation. The utility of the fractional residuals is not great. It is more informative to look at the residuals relative to the various flows and accumulations in the region.

A mass and energy balance with a small residual is necessary but not sufficient for an accurate numerical simulation. Because the system equations are a balance for each cell, and the method used for calculating the flows at specified-value boundary condition cells insures that the residual will be zero for those cells, and because the fluxes between the cells are conservative, errors in the global-balance equations will result from the following causes: (1) The approximate solution from use of the iterative-matrix equation solver; (2) the degree of convergence on density of the iteration on the

solution cycle of the three system equations; (3) the explicit treatment of the cross-dispersive flux terms; (4) the explicit and iterative treatment of well flow rates; and (5) roundoff error in special cases, such as wide variation in parameter magnitudes. Errors caused by discretization in time or space will not be revealed by these global-balance calculations. However, the

' inaccuracies resulting from too-long a time step under conditions of signifi-

" cantly nonlinear parameters will be evident. Significant nonlinearity can be caused by large variations in the density and viscosity fields, for example.

#### 4. COMPUTER CODE DESCRIPTION

### 4.1 CODE ORGANIZATION

The HST3D computer code is written in FORTRAN 77. Only code conforming to American National Standards Institute (ANSI) standards (American National Standards Institute, 1978) has been used to maximize program portability. The present version of the program code consists of a main routine and 49 subroutines. The program length is approximately 12,000 lines of code. Many FORTRAN statements occupy multiple lines. The following is a list of the routines and a brief description of their function:

HST3D - Main routine that drives the program execution. The basic steps are: (1) Read, ergor check, initialize, and write output for space allocation; (2) read, error check, initialize, and write output for static or time-invariant information; (3) read, error check, initialize and write output for transient information; (4) start the time-step calculations by calculating flow- and transport-equation coefficients, applying the boundary conditions, calculating the source-sink well terms; (5) perform the assembly and solution of the three equations in turn, iterating to convergence; (6) perform the summary calculations; (7) write the output information for the time plane; and (8) dump restart data to a disc file if desired. Then return to step 4 and continue until the time for a change in boundary conditions or source terms occurs. At this time return to step 3. Continue until the simulation is finished. Then (9), plot dependent variables as a function of time if desired; and (10) close files and terminate program execution.

The following subroutines are described in their order of execution:

READ1 - Reads the data pertaining to allocation of computer storage space for the problem.

ERROR1 - Checks for errors in the data read by READ1.

- INIT1 Initializes the spatial-allocation data, including the pointers for the two variable-partitioned arrays. If necessary, sets the inch-pound-to-metric conversion factors and their inverses, and sets the unit labels.
- WRITE1 Writes to a disc file information about the storage allocation and array-size requirements.
- READ2 Reads the data pertaining to all the time invariant or static information, including fluid and porous-media properties, grid geometry, equation-solution method, and desired output.

ERROR2 - Checks the data read by READ2 for errors.

- INIT2 Initializes the calculated static data for the simulation.
- WRITE2 Writes the static data to a disc file.
- READ3 Reads the transient data, including boundary condition and source-sink information, plus time-step, calculation, and printout information.

INIT3 - Initializes the calculated transient data.

ERROR3 - Checks the transient data read by READ3 for errors.

WRITE3 - Writes the transient data to a disc file.

COEFF - Calculates the coefficients for the flow-, heat- and solute-transport equations, and adjusts the automatic time step if selected. These coefficients include conductances, dispersion coefficients, and interstitial velocities.

WRITE4 - Writes the coefficients to a disc file as desired.

APLYBC - Applies the boundary conditions to the set of equations.

WELLSS - Calculates and applies the well source-sink terms.

ITER - Assembles and solves the three equations iteratively for a given time plane.

- SUMCAL Performs the summary calculations at the end of a time step. This includes flow, heat, and solute-mass balances, and flow rates.
- WRITE5 Writes to a disc file the desired information at the end of a time step. This may include pressure, temperature, and mass-fraction distributions, interstitial velocities, fluid viscosities, fluid densities, and summary tables of flow rates and balances of flow, heat, and solute mass.

DUMP - Dumps restart information to a disc file, if desired.

- PLOTOC Creates character-string plots of selected dependent variables (pressure temperature, or solute-mass fraction) as a function of time at the end of the simulation, and plots observed data if desired.
- CLOSE Closes files, deletes unused files, prints the total simulation time and the number of time planes, the number of restart and map records written, and any error messages.

The following subroutines are listed in approximate order of execution. Some are called from several routines, and some are optional.

ORDER - Determines the node numbering for the D4 reordering scheme for the direct-matrix equation solver.

- REWI Reads, error checks, echo writes, and initializes array elements that are input as zones of constant values over a rectangular prism of cells, or are input as node by node distributions.
- REWI3 The same as REWI, but for parameters that occur in sets of three, such as vectors or principal components of tensors.
- IREWI The same as REWI, but for parameters that are integers.
- VSINIT Calculates parameters for the viscosity function.
- ETOM1 Converts static data from inch-pound units to metric units.
- ETOM2 Converts transient data from inch-pound units to metric units.
- PRNTAR Writes one-dimensional or two-dimensional arrays in tabular form to a disc file.
- ZONPLT Creates two-dimensional maps on the printer of the porous-media zones contained in the simulation region.
- INTERP Performs one-dimensional or two-dimensional linear or bilinear tabular interpolation, as required.
- VISCOS Calculates fluid viscosity as a function of temperature and solutemass fraction.
- TOFEP Determines temperature as a function of enthalpy and pressure by tabular interpolation.
- WELRIS Performs the pressure and temperature calculation up or down the well-riser pipe, using simultaneous solution of the two ordinary differential equations.

ASEMBL - Assembles the coefficients of the modified flow, modified heat-transport, and solute-transport equations at each time step.

- CALCC Calculates the elements of the change in fluid-mass, change in heat, or change in solute-mass matrix for a given cell at each iteration at each time step.
- CRSDSP Calculates the components of the cross-dispersion tensor, that are evaluated explicitly in the transport equations at each iteration.
- D4DES Solves the matrix equation, using alternating-diagonal reordering and a direct Gaussian elimination algorithm.
- SOR2L Solves the matrix equation, using a two-line, successiveoverrelaxation algorithm, with the lines oriented in a selected coordinate direction.
- L2SOR Invokes the two-line, successive-overrelaxation solver for each coordinate direction to estimate the optimum overrelaxation parameter, and to solve the equations.
- MAP2D Creates two-dimensional contour maps on the printer of selected - variables with contour intervals divided into zones.

PLOT - Creates plots of pressure, temperature, or mass fraction versus time.

- ERRPRT Writes the error messages for a given simulation to a disc file.
- SBCFLO Calculates the flow rates at specified-value boundary-condition cells for the global balances.
- WBBAL Calculates flow rates for each well of fluid, heat, and solute for the summary calculations.

WBCFLO - Calculates the flow rates at the well-bore boundary for a single well in the cylindrical coordinate system.

- BSODE Integrates the coupled ordinary differential equations for pressure and temperature up or down the well-riser casing, using the Bulirsch-Stoer algorithm for rational polynomial extrapolation.
- WFDYDZ Calculates derivatives of pressure and temperature along the well-riser casing for the Bulirsch-Stoer integration algorithm.

If errors occur, the error checking that is in progress continues to completion, but then, information to that point in the calculations is written out, and the simulation is aborted.

A chart showing the main sequence of subroutine execution, the time step and transient data loops and the linkage between the subroutines appears in figure 4.1. The primary subroutines are on the left and the secondary and utility subroutines are to the right. The sequence of execution is from top to bottom of the leftmost column. Some utility subroutines are listed more than once for graphical clarity.

# 4.2 MEMORY ALLOCATION, ARRAY-SIZE REQUIREMENTS, AND SUBPROGRAM COMMUNICATION

A semi-dynamic method for array-storage allocation is employed in the simulator program (Akin and Stoddart, 1975, p. 114). Instead of dimensioning each of the arrays at some maximum value, a different approach is taken. The various arrays whose size depends on the number of nodes in the region, or the number of cells with a given type of boundary condition, or the number of wells, or the type of equation solver selected, are all contained in two large arrays; one array for real variables and one array for integer variables.

These two large arrays are partitioned into the required subarrays, based on the storage-allocation information provided. Pointer variables are used to indicate the location of the first element of each subarray in its large array. Thus, the variable-length subarrays are passed to the subroutines and functions though the calling arguments using the pointer variables. This

means that some subprograms have a large number of arguments. The advantage of this approach is that the lengths of the two large, variably partitioned arrays, VPA and IVPA, are set during compilation of the main routine, and only the main routine must be recompiled if the lengths are to be changed. The compiled length of the VPA array is contained in the variable ILVPA and the compiled length of the IVPA array is contained in the variable ILIVPA.

A rough estimate of the sizes required for the two large arrays is given by the following equations. For the large variably partitioned integer array,

ILI = 8 NXYZ + 6 NPMZ + 5 NWEL + NBC; (4.2.1) where

ILI is the estimated length of the large variably partitioned integer array;

NXYZ is the number of nodes in the region;

NPMZ is the number of porous medium zones;

NWEL is the number of wells; and

NBC is the number of boundary condition nodes.

For the large real array:

ILR = 70 NXYZ + 12 NPHZ + 60 NWEL + 60 NBC; (4.2.2)

where

ILR is the estimated length of the large variably partitioned integer array.

Equations 4.2.1 and 4.2.2 are based on solving all three equations and they overestimate the storage requirements in the interest of simplifying the estimate calculation. During the storage-allocation step, the actual required lengths of the large real and integer arrays are calculated and printed. Execution is aborted if insufficient space has been set during compilation. If redimensioning is necessary, the VPA and IVPA array sizes are redimensioned, and the variable ILVPA and is set to the compiled dimension of the VPA array, and the variable ILIVPA is set to the compiled dimension of the VPA array in the main HST3D program.

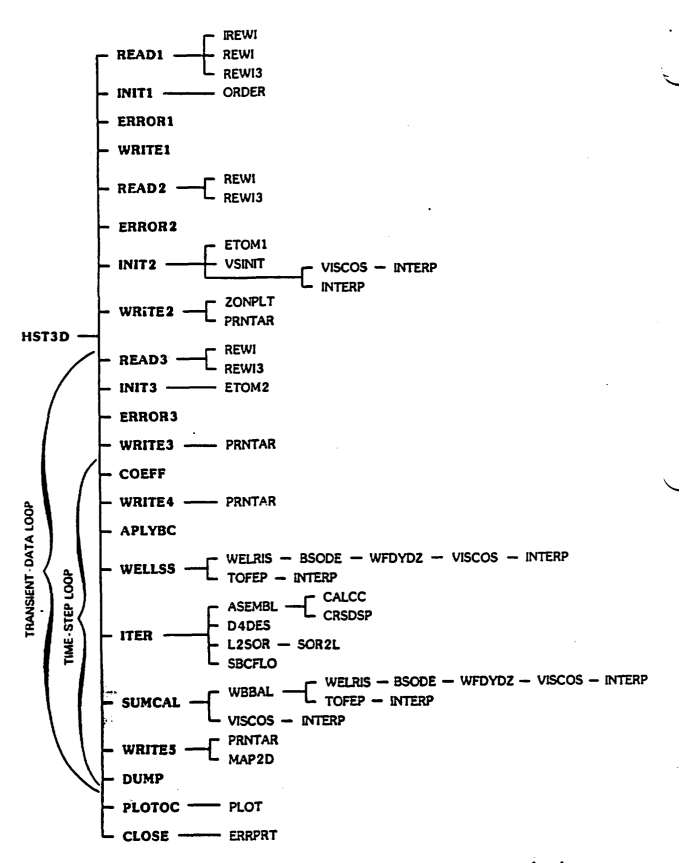


Figure 4.1.--Connection chart for the HST3D main program and subprogram showing routine hierarchy and calculation loops.

Some of the subarrays share storage space if they are used sequentially and the contents do not need to be retained for the duration of the simulation. Table 4.1 shows the partitioning of the large integer array; table 4.2 shows the partitioning of the large real array, with the shared storage indicated by subarrays on the same horizontal line. The size of each subarray and conditions for its inclusion (if optional) also are given. A significant savings in space is obtained by not including arrays that deal exclusively with either heat or solute transport in cases when transport of only one or the other is being simulated.

All other parameters and variables are passed through common blocks. Named common blocks are used, with the name being an abbreviation of the subprogram where that particular common block is first used with a suffix letter. There are many common blocks, because the variables are sorted into usage groups, so that each subprogram has access to only the common variables that it needs (as nearly as practical). All common blocks appear in the main program for easy reference and for static-storage allocation on some computer systems.

Another programming convention used is that, passing of subarrays through one subprogram to another is done by making the entire large array available to the calling subprogram along with the necessary pointer variables. This reduces the number of arguments for the calling subprogram.

		Other a	-	Conditi	ions for
		that			sion of
Subarray	Dimension	the	space	optional	subarrays
IBC	NXY2				
I1Z	NPMZ				••
12Z	NPMZ				
J1Z	NPMZ				
J2Z -	NPMZ				••
K1Z	NPMZ		•		••
K2Z	NPMZ				••
LPRNT	NXYZ	LEW, INZONE	, JWEL, IBCMAP		
IW	NWT			Wells	present
WL	NWT			Wells	present
LCTOPW	NWT			Wells	present
LCBOTW	NWT			Wells	present
WQMETH	NWT			Wells	present
MAIFC	NAIFC		,	Aquifer in b.c.	afluence function
MFBC	NFBC	**		Specif:	ied flux b.c.
MHCBC	NHCBC			Heat con	nduction b.c.
MLBC	NLBC			Leak	age b.c.
MSBC	NPTCBC			Specifie	d value b.c.
CI	6(NXYZ/2+1)			D4 mat	rix solver
IDIAG	NXYZ/2+1	**		D4 mat:	rix solver
RBW	NXYZ/2+1			D4 mat:	rix solver
ID4NO	NXYZ			D4 mat	rix solver

Table 4.1.--Space allocation within the large, variably partitioned, integer array, IVPA [--. no space sharing or no conditions; b.c., boundary condition]

a ....

	•	Other arrays that share	Conditions for inclusion of optional subarrays	
Subarray	Dimension	the space		
{	NX, NR	RR		
<b>x</b> {	NY		Cartesian coordinates	
	NZ			
M	NR-1		Cylindrical coordinates	
RX .	NXYZ	то		
RY	NXYZ		Cartesian coordinates	
RZ	NXYZ			
OROS	NPMZ		••	
BPM	NPMZ			
XX	NPMZ			
YY	NPMZ			
ZZ	NPMZ	<b></b>	**	
CPPM	NPMZ		Heat transport	
THX	NPMZ	••	Heat transport	
THY	NPMZ		Heat transport	
THZ	NPMZ		Heat transport	
LPHL	NPMZ		Heat or solute transport	
LPHT	NPMZ		Heat or solute transport	
BKD	NPMZ		Solute transport	
V .	NXYZ	POS		
MCV	NXYZ			
VK	NXYZ		Solute transport	
MCVK	NXYZ	<b></b>	Solute transport	
MHV	NXYZ	TOS	Heat transport	
MCHV	NXYZ		Heat transport	
	NXYZ	POW		
P	NXYZ	HWT, PNP		
	NXYZ	TOW	Heat transport	
T	NXYZ	UTBC, TNP, TQFLX	Heat transport	
	NXYZ	COW	Solute transport	
, C	NXYZ	UCBC, CNP, CQFLX	Solute transport	
)EN	NXYZ	TC		
IS	NXYZ	10		
H	NXYZ		Heat transport	
X	NXYZ			
Ŷ	NXYZ		Cartesian coordinates	
Z -	NXYZ	PCW	ATTEN CALATER	
TFX	NXYZ	HDPRNT, UPHILB	••	
FY	NXYZ	IDE MA DERTED	Cartesian coordinates	
TZ	NXYZ	INATEC IDENTE AMAD	Carcolan Containares	
HX		UVAIFC, UDENLB, AMAP	Heat transport	
	NXYZ	UDTHHC,QHFX,KTXPM	-	
IHXY	NXYZ		Heat transport, cartesia coordinates, full	

Table 4.2.--Space allocation within the large, variably partitioned real array, VPA [--, no space sharing or no conditions; b.c., mean boundary condition]

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dispersion tensor

real array, VPAContinued,			
		Other arrays	Conditions for
		that share	inclusion of
Subarray	Dimension	the space	optional subarrays
THXZ	NXYZ		Heat transport, full
			dispersion tensor
THY	NXYZ	QHFY , KTYPM	Heat transport, cartesian coordinates
THYX	NXYZ		Heat transport, full dispersion tensor, cartesian coordinates
THYZ	NXYZ		Heat transport, full dispersion tensor, cartesian coordinates
THZ	NXYZ	QHFZ, UKHCBC, KTZPM	Heat transport
THZX	NXYZ		Heat transport, full dispersion tensor
THZY	NXYZ	**	Heat transport, full dispersion tensor, cartesian coordinates
TSX	NXYZ	QSFX	Solute transport
TSXY	NXYZ		Solute transport, cartesian coordinates, full dispersion tensor
TSXZ	NXYZ	••	Solute transport, full dispersion tensor
TSY	NXYZ	QSFY	Solute transport, cartesian coordinates
TSYX	NXYZ		Solute transport, cartesian coordinates, full dispersion tensor
TSYZ	NXYZ		Solute transport, cartesian coordinates, full dispersion tensor
TSZ	NXYZ	QSFZ	Solute transport
TSZX	NXYZ	••	Solute transport, full dispersion tensor
TSZY	nxyz		Solute transport, full dispersion tensor, cartesian coordinates
SXX	NXYZ	QFFX,ARXBC,PCS	
syy	NXYZ	QFFY, ARYBC	Cartesian coordinates
SZZ	NXYZ	QFFZ, ARZBC	••
RF	NXYZ	APRNT, UVISLB	
RH	NXYZ		Heat transport
RH1	NXYZ		Heat transport
RS	NXYZ	CCW	Solute transport
RS1	NXYZ	**	Solute transport
URR1	NXYZ		Solute transport

Table 4.2.--Space allocation within the large, variably partitioned real array, VPA--Continued.

Other arrays			Conditions for	
· .		that share	inclusion of	
Subarray	Dimension	the space	optional subarrays	
. ,				
RHS	NXYZ	FRAC, UZELB, TCS	Solute transport	
C24	NXYZ	UBBLB	· •••	
CC34	NXYZ			
CC35	NXYZ	TCW		
78.	7 •NXYZ	VXX,AA1,MOBW,UDENBC,UKLB	•=	
		VYY,AA2		
	· · · · ·	VZZ,AA3	••	
	an a film	AA4		
<b>VI</b>	- NWEL·NZ		Wells present	
QWLYR	NWEL·NZ	WCF	Wells present	
QHLYR	NWEL · NZ		Wells present, heat	
			transport	
QSLYR	NWEL·NZ		Wells present, solute	
			transport	
DQWDPL	NWEL • NZ		Wells present	
WRANGL	NWEL		Wells present	
WBOD	NWEL		Wells present	
	NWEL			
WRISL	NWEL		Wells present	
WRID			Wells present	
WRRUF	NWEL		Wells present	
KTHWR	NWEL	••	Wells present, heat transport	
DTHWR	NWEL		Wells present, heat	
		• •	transport	
KTHAWR	NWEL	••	Wells present, heat	
		•	transport	
HTCWR	NWEL	••	Wells present, heat	
	•		transport	
TABWR	NWEL	••	Wells present, heat	
:	<u>.</u>		transport	
TATWR	NWEL		Wells present, heat	
· · ·		· .	transport	
TWKT	NWEL		Wells present, heat transport	
TWSUR	NWEL	•• *	Wells present, heat	
	67 W 66		transport	
EHWKT	NWEL	<b></b>	Wells present, heat	
LIIWA I	NWLL .		transport	
FULKIE	17.7E T	-	Wells present, heat	
EHWSUR	NWEL		-	
DIRTE			transport Valle present	
PWKTS	NWEL		Wells present	
PWKT	NWEL		Wells present	
DPWKT	NWEL	<b>▲●</b>	Wells present	
PWSURS	NWEL	•• * *	Wells present	
PWSUR	NWEL		Wells present	

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Table 4.2.--Space allocation within the large, variably partitioned real array, VPA--Continued

CWKT       NWEL        Wells present, solute transport         QWV       NWEL        Wells present         QWM       NWEL        Wells present         QHW       NWEL        Wells present, heat         QSW       NWEL        Wells present, heat         QSW       NWEL        Wells present, solute         QSW       NWEL        Wells present, solute         QSW       NWEL        Wells present, heat         QSW       NWEL        Wells present, solute         QSW       NWEL        Wells present, heat         QSW       NWEL        Wells present, heat         WFICUM       NWEL        Wells present, heat         WHICUM       NWEL        Wells present, solute         WHPCUM       NWEL        Wells present, solute         WHY NWEL        Wells present, solute       transport         WSPCUM       NWEL        Wells present, solute       transport         VASBC       7-NPTCBC        Specified-value b.c.       RHSBC         VASBC       NPTCBC			Other arrays that share	Conditions for inclusion of
UNVNWELWells presentQHMNWELWells present,QHWNWELWells present,QSWNWELWells present,QSWNWELWells present,QSWNWELWells present,WFICUMNWELWells present,WFPCUMNWELWells present,WHICUMNWELWells present,WHOUMNWELWells present,WSICUMNWELWells present,NNELWells present,WSICUMNWELWESCYessent,NUELWells present,VASBC7-NPTCBCRHSBCNPTCBCSpecified-value b.c.Resent,QFSBCNPTCBCSpecified-value b.c.andheat transportSpecified-value b.c.QSSBCNPTCBCSpecified-value b.c.andcSBCNPTCBCSpecified-value b.c.andsolute transportSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.cGFECNFBCSpecified-flux b.cGRFBCNFBCSpecified-flux b.c.andsolute transportSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.cGFEXNFBC-	Subarray	Dimension	the space	optional subarrays
DVVNWELWells presentDMWNWELWells presentDNWNWELWells presentDSWNWELWells present, heatDSWNWELWells present, soluteDSWNWELWells presentWFICUMNWELWells presentWHICUMNWELWells present, heatWHICUMNWELWells present, heatWHPCUMNWELWells present, soluteWSICUMNWELWells present, soluteWSPCUMNWELWells present, soluteVASBC7.NPTCBCSpecified-value b.c.RISBCNFTCBCSpecified-value b.c.QSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-value b.c.MRSBCNFTCBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecifie	CWKT	NWEL		
QMMNWELWells presentQNWNWELWells present, heatQSWNWELWells present, solutewFPCUMNWELWells presentWFPCUMNWELWells presentWHCUMNWELWells presentWHCUMNWELWells present, heatWHCUMNWELWells present, heatWHCUMNWELWells present, soluteWSPCUMNWELWells present, soluteWSPCUMNWELSpecified-value b.c.WSPCUMNWELSpecified-value b.c.WSPCUMNWELSpecified-value b.c.WSPCUMNWELSpecified-value b.c.QFSBCNFTCBCSpecified-value b.c.MSSGNFTCBCSpecified-flux b.c.QSSBCNFTCBCSpecified-flux b.c.QSSBCNFTCBC </td <td>0WV</td> <td>NWEL</td> <td></td> <td></td>	0WV	NWEL		
NWELWells present, heat transportQSWNWELWells present, solute transportWFICUMNWELWells presentWFICUMNWELWells presentWHICUMNWELWells present, heat transportWHPCUMNWELWells present, heat transportWRCUMNWELWells present, solute transportWSICUMNWELWells present, solute transportWSPCUMNWELWells present, solute 	•			
QSWNWELWells present, solute transportWFICUMNWELWells presentWFPCUMNWELWells present, heat transportWHICUMNWELWells present, heat transportWHUMNWELWells present, solute transportWSICUMNWELWells present, solute transportWSICUMNWELWells present, solute transportVASBC7 NPTCBCSpecified-value b.c.VASBC7 NPTCBCSpecified-value b.c.QFSBCNPTCBCSpecified-value b.c.QFSBCNPTCBCSpecified-value b.c.QFSBCNPTCBCSpecified-value b.c.QSSBCNPTCBCSpecified-value b.c.QSSBCNPTCBCSpecified-value b.c.QSSBCNPTCBCSpecified-value b.c.QSSBCNPTCBCSpecified-value b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.QFFBCNFBCSpecified-flux b.c.QFFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXFBCNFBCSpecified-flux b.c.ARXF	•		••	
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	CFLX	NFBC		Specified-flux b.c. and
	ALBC	NLBC		

Table 4.2.--Space allocation within the large, variably partitioned real array, VPA--Continued

Subarran	Dimension	Other arrays that share	Conditions for inclusion of
Subarray	Dimension	the space	optional subarrays
BLBC	NLBC	••	Leakage b.c.
KLBC	NLBC	· ••	Leakage b.c.
ZELBC	NLBC		Leakage b.c.
BBLBC	NLBC		Leakage b.c.
HLBC	NLBC	PHILBC	Leakage b.c.
DENLBC	NLBC		Leakage b.c.
VISLBC	NLBC		Leakage b.c.
TLBC	NLBC		Leakage b.c. and heat
CLBC	NLBC	••	transport Leakage b.c. and solute
			transport
QFLBC	NLBC		Leakage b.c.
QHLBC	NLBC		Leakage b.c. and heat transport
QSLBC	NLBC	· ·	Leakage b.c. and solute transport
AAIF	NAIFC		Aquifer-influence-function
BAIF	NAIFC		Aquifer-influence-function b.c.
VAIF	NAIFC		Aquifer-influence-function
WCAIF	NAIFC		Aquifer-influence-function
DENOAR	NAIFC		b.c. Aquifer-influence-functio
PAIF	NAIFC		b.c. Aquifer-influence-function
TAIF	NAIFC	•• × · · · · · · · · · · · · · · · · · · ·	b.c. Aquifer-influence-function
CAIF	NAIFC	••	b.c. and heat transport Aquifer-influence-function
		•	b.c. and solute
QFAIF	NAIFC	••	transport Aquifer-influence-function
QHAIF	NAIFC	•• • • •	b.c. Aquifer-influence-function
4.0.22	511142 U		b.c. and heat transport
QSAIF	NAIFC	••	Aquifer-influence-function b.c. and solute transport
ZHCBC	NHCN		Heat-conduction b.c.
A1HC	NHCN	••	Heat-conduction b.c.
A2HC	NHCN		Heat-conduction b.c.
A3HC	NHCN		Heat-conduction b.c.

Table 4.2.--Space allocation within the large, variably partitioned real array, VPA--Continued

}

Subarray	Dimension	Other arrays that share the space	Conditions for inclusion of optional subarrays
DTHHC QHCBC DQHCDT THCBC TPHCBC A4	NHCBC NHCBC NHCBC • NHCN NHCBC • NHCN NHCBC • NHCN MAXNAL (D4 SOLVER) OR 7 • NXYZ (L2SOR SOLVER)	  D TM1 TM2 TP1 TP2 XX XXN	Heat-conduction b.c. Heat-conduction b.c. Heat-conduction b.c. Heat-conduction b.c. Heat-conduction b.c. D4 direct solution method or two-line, successive overrelaxation iterative-solution method

Table 4.2.--Space allocation within the large, variably partitioned real array, VPA--Continued

# 4.3 FILE USAGE

The heat- and solute-transport program uses several sequential-access files on disc. The following list gives the FORTRAN unit number and a description of each file:

FILE 5 Input data with comments stripped out.

FILE 6 Output data in character form for the line printer or video screen.

FILE 7 Plot data for dependent variables versus time.

FILE 8 Output data for a restart run.

FILE 9 Input data for a restart run.

FILE 10 Input data with comment lines

FILE 11 Echo of the input data as it is read.

These files must be declared or established on the computer system before program execution. The present version of the program also requires preopening of these files before starting execution. File names are assigned when the files are opened, using the job-control language of the computer employed.

# 4.4 INITIALIZATION OF VARIABLES

In general, all real and integer variables are set to zero and logical variables are set to false by explicitly coded instructions at the beginning of program execution. Certain input parameters are set to default values if no data or data values of zero are input. These are indicated in the inputfile description. Two tables are initialized for the enthalpy of pure water as a function of temperature and pressure. These tables are used for the heat-transport calculation.

# 4.5 PROGRAM EXECUTION

The program is designed to be executed by the job-control language (JCL) or command-procedure language (CPL) of the computer employed. The data-input file for a given program must have been created previously. The job-control language statements should declare and open the FORTRAN files listed above, assign file names, then start program execution. At the end of the simulation, the JCL or CPL statements should close all files and delete scratch files. Unused files are deleted by the program in the CLOSE subroutine. Execution of the command-procedure, control-statement file is usually done interactively at a terminal, so file names can be provided by the user.

# 4.6 RESTART OPTION

The program has the option for restarting from an intermediate or ending time of the simulation. The user may specify that restart information is to be written to the restart file every nth time step (record 3.8.3 of the input file). Then, the ability to continue in a later run is created. This is useful; for example, if boundary conditions or sources need to be changed, based on the results of the simulation, or if a steady-state flow field needs to be established, before a heat or solute source is introduced. The other use of the restart option is to specify that restart information is to be written at periodic time-step intervals. This is insurance against computersystem failure, so that long simulation runs will not have to be repeated entirely.

The information written to disc for restart consists of the two large, variably partitioned arrays, and those common blocks that contain parameters and variables that are active during the simulation time-step loop. This information can be written only at the end of a time step. To restart the simulation, a small amount of data is needed in the input file, including the first three lines of the READ1 record group and the transient-boundary condition and fluid-source information of the READ3 record group. After the restart-data file is read, execution continues at the beginning of the subroutine (READ3) that reads the transient information, shown in figure 4.1. Only the transient data that are to be changed from the last values of the previous simulation need to be read at the beginning of a restart. From this point onward in the simulation, there is no difference between a restart run and one that has just gone through a change in transient-boundary conditions. In particular, time is measured relative to the beginning of the original simulation period.

One consequence of the way restarts are implemented is the restriction that the number and type of boundary condition and source nodes cannot be changed from the values for the original simulation run. To do so will cause a fatal error resulting from a file length conflict. At the beginning of a restart, the program will attempt to find the restart file whose time-plane value agrees within 0.1 percent of the specified time value for restarting. Failure to locate the specified restart file causes a fatal error.

# 5. THE DATA-INPUT FILE

. . .

The data-input file has two general characteristics: (1) It is free format for ease of preparation at a computer terminal; and (2) it may be freely commented for rapid identification of the data items. The free format is supported by FORTRAN 77, and is sometimes referred to as list-directed input.

## 5.1. LIST DIRECTED INPUT

The data values may be located in any position of the record, provided they correspond in number and type with the input list. Data are separated by commas or blanks, with multiple blanks allowed. Character strings must be enclosed in quotation marks (apostrophies on some computers). There is a third delimiter in addition to a comma and a blank: the slash, /. A slash terminates a record and any remaining items in the input list are left unchanged from their previous values. On some computer systems, there must be a space before the slash. A data item within an input list may be left unchanged by separating the preceding item from the subsequent item with two commas; in other words, making no entry for that item.

The list-directed input of a record continues until the end-of-record slash is encountered, or the input list is satisfied. If the input list is not satisfied at the end of a data line, the program will continue to read additional data lines, until the list is satisfied, or the end of the file is reached. Having an insufficient number of data items in the input file is a common error. It usually results from a misinterpretation of the amount or type of data required by a given program option.

List-directed input also allows for a repeat count. Data in the form:

n\*d;

(5.1.1)

where

n is the repeat count integer; and d is the data item;

causes n consecutive values of d to be input.

# 5.2. PREPARING THE DATA-INPUT FILE

To simplify the preparation of the data-input file, a file is available containing only comments. This file is presented as table 5.1. These comments identify all the input-variable names, indicate the logical ones, show the conditions for the optional items, and give the default values, where used. The user actually can enter the data between the lines of a copy of this file, using section 5.2.2 as a guide. This data-input form should make it easier to create and modify the data-input file. Only the variable names used in the program are given. For definitions see section 5.2.2 or 11.1. Optional input records are indicated by (O) followed by the logical variables that must be true, or the numerical conditions that must be met for inclusion of a given input record. The numbers in brackets after a variable give the record where that variable is read. The following section contains a line-by-line presentation of the data-file form with an explanation of all the variables and options.

Many published sources exist of fluid and porous-medium properties, and transport parameters such as compressibilities, heat capacities, and equilibrium distribution coefficients. Typical examples are Perry and others (1963), Clark (1966), Weast and others (1964), and Mercer and others (1982).

#### 5.2.1. General Information

Comment lines are numbered in the format C.N1.N2.N3 where C denotes a comment record; N1 is the read-group number; and N2.N3 is the record or line number. The read-group number denotes which subroutine, READ1, READ2, READ3,

or PLOTOC, reads that record for the values 1 through 4 respectively. The record number identifies a line of input data, with the two component number enabling a logical group structure to be assigned. A suffix letter associated with N2 or N3 indicates that one of the numbered set with that letter must be selected exclusively. Optional data records are indicated to the right of the record list of variables, and the conditions under which these data are required are explained. If a particular input record is not needed, it is manditory that it be omitted. A record number in square brackets following a variable indicates the record where that variable is first set.

Input by i,j,k range means that an array of data is to be specified for a rectangular prism of nodes. The extent of this prism is defined by the ranges of the i,j, and k indices. One may specify a plane, or a line, or a point instead of a prism by making the appropriate indices equal. The value to be entered into the array locations pertaining to the nodes contained with the prism is specified next. There is the capability to replace, multiply or add to existing data within the prism. The specified value fills the specified array locations or operates on the existing values in those locations. In some cases, three arrays are affected for the nodes within the specified prism. There is a fourth option for data input which allows for a sequence of different values to be entered into consecutive node locations within the prism. The data will be loaded in order of increasing node number, that is, in order of increasing i, then increasing j, and then increasing k. With this input scheme, subregions of uniform parameters within the simulation region are easily defined. A parameter distribution that varies continuously in space is less convenient to define. It is permissible to use rectangular prisms that include cells that are outside the simulation region.

Table 5.1.--Data-input form

C....HST DATA-INPUT FORM NOTES: C... INPUT LINES ARE DENOTED BY C.N1.N2.N3 WHERE C... N1 IS THE READ GROUP NUMBER, N2.N3 IS THE RECORD NUMBER C... A LETTER INDICATES AN EXCLUSIVE RECORD SELECTION MUST BE WADE C... I.E. A OR B OR C C... **c**... (0) - OPTIONAL DATA WITH CONDITIONS FOR REQUIREMENT A RECORD NUMBER IN SQUARE BRACKETS IS THE RECORD WHERE THAT PARAMETER IS FIRST SET C... C..... INPUT BY I.J.K RANGE FORMAT IS: C.O.1.. 11.12.J1.J2.K1.K2 C.O.2.. VAR1, INOD1, [VAR2, INOD2, VAR3, INOD3] C... USE AS MANY OF LINE 0.1 & 0.2 SETS AS NECESSARY C... END WITH LINE 0.3 C.O.3.. 0 / THE SPACE IS REQUIRED C... (NNN) - INDICATES THAT THE DEFAULT NUMBER, NNN, IS USED IF A ZERO IS ENTERED FOR THAT VARIABLE C... (T/F) - INDICATES A LOGICAL VARIABLE C... (I) - INDICATES AN INTEGER VARIABLE ſ\_\_\_\_\_ C.....START OF THE DATA FILE C.....DIMENSIONING DATA - READ1 C.1.1 .. TITLE LINE 1 C.1.2 .. TITLE LINE 2 C.1.3 .. RESTRT(T/F), TIMRST C.... IF RESTRT IS TRUE, PROCEED TO READ3 GROUP [3.1] C.1.4 .. HEAT.SOLUTE.EEUNIT.CYLIND.SCALNF: ALL (T/F) C.1.5 .. NX.NY.NZ.NHCN C.1.6 .. NPTCBC, NFBC, NAIFC, NLBC, NHCBC, NWEL C.1.7 .. NPHZ C.1.B .. SLMETH[I], LCROSD(T/F) C.1.9 .. IBC BY I, J, K RANGE (0.1-0.3) , WITH NO IMOD PARAMETER, FOR EXCLUDED CELLS C.1.10 .. RDECHO(T/F) C-----C....STATIC DATA - READ2 C....OUTPUT INFORMATION C.2.1 .. PRTRE(T/F) C....COORDINATE GEOMETRY INFORMATION C..... RECTANGULAR COORDINATES C.2.2A.1 .. UNIGRX, UNIGRY, UNIGRZ; ALL (T/F); (0) - NOT CYLIND [1.4] C.2.2A.2A .. X(1),X(NX);(0) - UNIGRX [2.2A.1] C.2.2A.2B .. X(I);(0) - NOT UNIGRX [2.2A.1] C.2.2A.3A .. Y(1),Y(NY);(0) - UNIGRY [2.2A.1] C.2.2A.38 .. Y(J);(O) - NOT UNIGRY [2.2A.1] C.2.2A.4A .. Z(1), Z(NZ); (0) - UNIGRZ [2.2A.1] C.2.2A.4B .. Z(K);(0) - NOT UNIGRZ [2.2A.1] 1 C.... CY INDRICAL COORDINATES

#### Table 5.1.--Data-input form--Continued

C.2.28.1A .. R(1),R(NR),ARGRID(T/F);(0) - CYLIND [1.4] C.2.2B.1B .. R(I);(0) - NOT ARGRID [2.2B.1A];(0) - CYLIND [1.4] C.2.28.2 .. UNIGRZ(T/F):(0) - CYLIND [1.4] C.2.2B.3A .. Z(1), Z(NZ); (0) - UNIGRZ [2.2B.3A], CYLIND [1.4] C.2.2B.3B .. Z(K);(0) - NOT UNIGRZ [2.2B.3A],CYLIND [1.4] C.2.3.1 .. TILT(T/F);(0) - NOT CYLIND [1.4] C.2.3.2 .. THETXZ, THETYZ, THETZZ; (0) - TILT [2.3.1] AND NOT CYLIND [1.4] C....FLUID PROPERTY INFORMATION C.2.4.1 .. 8P C.2.4.2 .. PO, TO, WO, DENFO C.2.4.3 .. W1, DENF1:(0) - SOLUTE [1.4] C.2.5.1 .. NOIVO, TVFO(1), VISTFO(1), I=1 TO NOTVO; (0) - HEAT [1.4] OR HEAT [1.4] AND SOLUTE [1.4] OR .NOT.HEAT AND .NOT.SOLUTE [1.4] C.2.5.2 .. NOTV1,TVF1(I),VISTF1(I),I=1 TO NOTV1;(0) - SOLUTE [1.4] AND HEAT [1.4] C.2.5.3 .. NOCV, TRVIS, CVIS(1), VISCTR(1), I=1 TO NOCV; (0) - SOLUTE [1.4] C.....REFERENCE CONDITION INFORMATION C.2.6.1 .. PAATH C.2.6.2 .. POH, TOH C.....FLUID THERMAL PROPERTY INFORMATION C.2.7 .. CPF, KTHF, BT; (0) - HEAT [1.4] C.....SOLUTE INFORMATION C.2.8 .. DM, DECLAM; (0) - SOLUTE [1.4] C....POROUS MEDIA ZONE INFORMATION C.2.9.1 .. IPNZ\_[12(IPNZ)\_12Z(IPNZ)\_J1Z(IPNZ)\_J2Z(IPNZ)\_K1Z(IPNZ)\_K2Z(IPNZ) C..... USE AS MANY 2.9.1 LINES AS NECESSARY C.2.9.2 .. END WITH 0 / C.....POROUS NEDIA PROPERTY INFORMATION C.2.10.1 .. KXX(1PHZ),KYY(1PHZ),KZZ(1PHZ),1PHZ=1 TO NPHZ [1.7] C.2.10.2 .. POROS(IPMZ), IPMZ=1 TO NPMZ [1.7] C.2.10.3 .. ABPH(IPMZ), IPMZ=1 TO NPMZ [1.7] C.....POROUS MEDIA THERMAL PROPERTY INFORMATION C.2.11.1 .. RCPPH(IPMZ), IPMZ=1 TO MPMZ [1.7]; (0) - HEAT [1.4] C.2.11.2 .. KTXPW(IPMZ),KTYPM(IPMZ),KTZPM(IPMZ),IPMZ=1 TO NPMZ [1.7];(0) - HEAT [1.4] C.....POROUS MEDIA SOLUTE AND THERMAL DISPERSION INFORMATION C.2.12 .. ALPHL(IPMZ).ALPHT(IPMZ), IPMZ=1 TO NPMZ [1.7];(0) - SOLUTE [1.4] OR HEAT [1.4] C.....POROUS MEDIA SOLUTE PROPERTY INFORMATION C.2.13 .. DBKD(IPHZ), IPHZ=1 TO NPHZ [1.7];(0) - SOLUTE [1.4] C.....SOURCE-SINK WELL INFORMATION C.2.14.1..RDWDEF(T/F);(0) - HWEL [1.6] > 0C.2.14.2... IMPOW(T/F);(0) - NWEL [1.6] > 0 AND NOT CYLIND [1.4] C.2.14.3. .. INEL, IN, JW, LCBOTW, LCTOPW, WBOD, WOMETH[1]; (0) - ROWDEF [2.14.1]. C.2.14.4 .. WCF(L); L = 1 TO NZ (EXCLUSIVE) BY ELEMENT C.2.14.5 .. WRISL, WRID, WRRUF, WRANGL; (0) - ROWDEF [2.14.1] AND WRCALC(WOMETH [2.14.3] >30) C.2.14.6 .. HTCWR.DTHAWR.KTHAWR,KTHWR, TABWR, TATWR; (0) - RDWDEF [2.14.1] WRCALC(WOMETH [2.14.3] >30) AND HEAT [1.4] C.... USE AS MANY 2.14.3-6 LINES AS NECESSARY C.2.14.7 .. END WITH 0 / 1

```
C.2.14.8 .. MXITQW[14],TOLDPW[6.E-3],TOLFPW[.001],TOLQW[.001],DAMWRC[2.],DZMIN(.01),EPSWR[.001];(0) - RDWDEF [2.14.1]
C....
                AND WRCALC(WOMETH 2.14.3] >30)
C....BOUNDARY CONDITION INFORMATION
C....
           SPECIFIED VALUE B.C.
C.2.15 .. IBC BY I.J.K RANGE (0.1-0.3) WITH NO INOD PARAMETER, :(0) - NPTCBC [1.6] > 0
           SPECIFIED FLUX B.C.
C....
C.2.16 .. IBC BY I.J.K RANGE (0.1-0.3) WITH NO INOD PARAMETER;;(0) - NFBC [1.6] > 0
           AQUIFER AND RIVER LEAKAGE B.C.
C....
C.2.17.1 .. IBC BY I.J.K RANGE (0.1-0.3) WITH NO INOD PARAMETER;(0) - NLBC [1.6] > 0
C.2.17.2 .. KLBC,BBLBC,ZELBC BY I,J,K RANGE (0.1-0.3);(0) - NLBC [1.6] > 0
                RIVER LEAKAGE B.C.
C....
C.2.17.3 .. I1, I2, J1, J2, KRBC, BBRBC, ZERBC; (0) - HLBC [1.6] > 0
C.2.17.4 .. END WITH 0 /
C....
           AQUIFER INFLUENCE FUNCTIONS
C.2.18.1 .. IBC BY I.J.K RANGE (0.1-0.3) WITH NO INOD PARAMETER;(0) - NAIFC [1.6] >0
C.2.18.2 .. UVAIFC BY I.J.K RANGE {0.1-0.3};(0) - NAIFC [1.6] > 0
C.2.18.3...IAIF;(0) - HAIFC [1.6] > 0
C.....TRANSIENT, CARTER-TRACY A.I.F.
C.2.18.4 .. KOAR.ABOAR.VISOAR.POROAR.BOAR.RIOAR.ANGOAR;(0) - IAIF [2.18.3] = 2
           HEAT CONDUCTION B.C.
C....
C.2.19.1 .. IBC BY I.J.K RANGE (0.1-0.3) , WITH NO IMOD PARAMETER .FOR HCBC NODES:(0) - HEAT (1.4) AND NHCBC (1.6) > 0
C.2.19.2 .. ZHCBC(K);(O) - HEAT [1.4] AND NHCBC [1.6] > 0
C.2.19.3 .. UDTHHC BY I, J, K RANGE (0.1-0.3) FOR HCBC HODES: (0) - HEAT-[1.4] AND NHCBC [1.6] > 0
C.2.19.4 .. UKHCBC BY I, J, K RANGE (0.1-0.3) FOR HCBC NODES; (0) - HEAT [1.4] AND NHCBC [1.6] >0
C....FREE SURFACE B.C.
C.2.20 .. FRESUR(T/F).PRTCCH(T/F)
C....INITIAL CONDITION INFORMATION
C.2.21.1 .. ICHYDP.ICT.ICC: ALL (T/F); IF NOT.HEAT, ICT = F, IF NOT.SOLUTE, ICC = F
C.2.21.2 .. ICHWT(T/F);(0) - FRESUR [2.20]
C.2.21.3A .. ZPINIT, PINIT; (0) - ICHYDP [2.21.1] AND NOT ICHWT [2.21.2]
C.2.21.3B .. P BY 1, J.K RANGE (0.1-0.3); (0) - NOT ICHYDP (2.21.1) AND NOT ICHWT (2.21.2)
C.2.21.3C .. HWT BY I, J, K RANGE (0.1-0.3); (0) - FRESUR [2.20] AND ICHWT [2.21.2]
C.2.21.4A .. NZTPRO,ZT(I),TVD(I),I=1,NZTPRO;(O) - HEAT [1.4] AND NOT ICT [2.21.1], LIHIT OF 10
C.2.21.4B .. T BY 1,J,K RANGE (0.1-0.3);(0) - HEAT [1.4] AND ICT [2.21.1]
C.2.21.5 .. NZTPHC, ZIHC(1), TV2HC(1); (0) - HEAT [1.4] AND NHCBC [1.6] > 0, LIMIT OF 5
C.2.21.6 .. C BY I, J, K RANGE (0.1-0.3);(0) - SOLUTE [1.4] AND ICC [2.21.1]
C.....CALCULATION INFORMATION
C.2.22.1 .. FDSNTH.FDTNTH
C.2.22.2 .. TOLDEN(.001).MAXITN(5)
C.2.22.3 .. NTSOPT(5), EPSSOR(.00001), EPSONG(.2), MAXIT1(50), MAXIT2(100); (0) - SLMETH (1.8) = 2
C....OUTPUT INFORMATION
C.2.23.1 .. PRTPMP_PRTFP_PRTIC_PRTBC_PRTSLM_PRTWEL: ALL (T/F)
C.2.23.2... IPRPTC, PRTDV(T/F); (0) - PRTIC [2.23.1]
C.2.23.3 .. ORENPR[I];(0) - NOT CYLIND [1.4]
C.2.23.4 .. PLTZON(T/F);(0) - PRTPMP [2.23.1]
```

Table 5.1.--Data-input form--Continued

C.2.23.5 .. OCPLOT(T/F) C------C..... TRANSIENT DATA - READS C.3.1 .. THRU(T/F) C.....IF THRU IS TRUE PROCEED TO RECORD 3.99 C....THE FOLLOWING IS FOR NOT THRU C.....SOURCE-SINK WELL INFORMATION  $C.3.2.1..RDWFLO(T/F)_RDWHD(T/F):(0) - HWEL [1.6] > 0$ C.3.2.2 .. IWEL, ONV, PWSUR, PWKT, TWSRKT, CWKT; (0) - RDWFLO [3.2.1] OR RDWHD [3.2.1] C....USE AS MANY 3.2.2 LINES AS NECESSARY C.3.2.3 .. END WITH 0 / C.....BOUNDARY CONDITION INFORMATION SPECIFIED VALUE B.C. C.... C.3.3.1 ... RDSPBC, RDSTBC, RDSCBC, ALL(T/F); (0) - NOT CYLIND [1.4] AND NPTCBC [1.6] > 0 C.3.3.2 .. PNP B.C. BY I.J.K RANGE (0.1-0.3):(0) - RDSPBC [3.3.1] C.3.3.3 .. TSBC BY I, J, K RANGE (0.1-0.3); (0) - RDSPBC [3.3.1] AND HEAT [1.4] C.3.3.4 .. CSBC BY I.J.K RANGE (0.1-0.3); (0) - RDSPBC [3.3.1] AND SOLUTE [1.4] C.3.3.5 .. THP B.C. BY I,J,K RANGE (0.1-0.3):(0) - RDSTBC [3.3.1] AND HEAT [1.4] C.3.3.6 .. CMP B.C. BY I, J, K RANGE (0.1-0.3); (0) - RDSCBC [3.3.1] AND SOLUTE [1.4] C.... SPECIFIED FLUX C.3.4.1 .. RDFLXQ, RDFLXH, RDFLXS, ALL(T/F); (0) - NFBC [1.6] > 0 C.3.4.2 .. QFFX,QFFY,QFFZ B.C. BY I,J,K RANGE (0.1-0.3);(0) - RDFLXQ [3.4.1] C.3.4.3 .. UDENBC BY I, J, K RANGE (0.1-0.3); (0) - RDFLX0 [3.4.1] C.3.4.4 .. TFLX B.C. BY I,J,K RANGE {0.1-0.3};(0) - RDFLXQ [3.4.1] AND HEAT [1.4] C.3.4.5 .. CFLX B.C. BY I.J.K RANGE (0.1-0.3);(0) - RDFLX0 [3.4.1] AND SOLUTE [1.4] C.3.4.6 .. QHFX.QHFY.QHFZ B.C. BY I.J.K RANGE (0.1-0.3):(0) - RDFLXH [3.4.5] C.3.4.7 .. QSFX,QSFY,QSFZ B.C. BY I,J,K RANGE (0.1-0.3);(0) - RDFLXS [3.4.1] LEAKAGE BOUNDARY C.... C.3.5.1..ROLBC(T/F):(0) - MLBC[1.6] > 0C.3.5.2 .. PHILBC, DEMLBC, VISLBC BY I, J,K RANGE (0.1-0.3);(0) - RDLBC [3.5.1] C.3.5.3 .. TLBC BY I.J.K RANGE (0.1-0.3):(0) - RDLBC [3.5.1] AND HEAT [1.4] C.3.5.4 .. CLBC BY I.J.K RANGE (0.1-0.3):(0) - ROLBC [3.5.1] AND SOLUTE [1.4] RIVER LEAKAGE C.... C.3.5.5 .. I1,I2,J1,J2,HRBC,DENRBC,VISRBC,TRBC,CRBC;(0) - RDLBC [3.5.1] C....USE AS MANY 3.5.5 LINES AS HECESSARY C.3.5.6 .. END WITH 0 / A.I.F. B.C. C.... C.3.6.1..RDAIF(T/F); (0) - MAIFC [1.6] > 0C.3.6.2 .. DEMOAR BY I, J, K RANGE (0.1-0.3);(0) - RDAIF [3.6.1] C.3.6.3 .. TAIF BY I, J, K RANGE (0.1-0.3); (0) - RDAIF (3.6.1) AND HEAT [1.4] C.3.6.4 .. CAIF BY I.J.K RANGE (0.1-0.3):(0) - RDAIF [3.6.1] AND SOLUTE [1.4] C....CALCULATION INFORMATION C.3.7.1 .. RDCALC(T/F) C.3.7.2 .. AUTOTS(T/F);(0) - RDCALC [3.7.1] C.3.7.3.A .. DELTIM; (0) - RDCALC [3.7.1] AND NOT AUTOTS [3.7.2]

#### Table 5.1.--Data-input form--Continued

```
C.3.7.3.B .. DPTAS(5E4), DTTAS(5.), DCTAS(.25), DTINNN(1.E4), DTINNX(1.E7); (0) - RDCALC [3.7.1] AND AUTOTS [3.7.2]
C.3.7.4 .. TINCHG
C....OUTPUT INFORMATION
C.3.8.1 .. PRIVEL_PRIDV.PRISLM_PRIKD_PRIPTC.PRIGF8.PRIWEL,PRIBCF; ALL []
C.3.8.2 .. IPRPTC; (0) - IF PRIPTC [3.8.1] NOT = 0
C.3.8.3 .. CHKPTD(T/F), NTSCHK, SAVLDO(T/F)
C....CONTOUR MAP INFORMATION
C.3.9.1 .. RDHPDT, PRTMPD; ALL (T/F)
C.3.9.2 .. HAPPTC.PRIMAP[1]:(0) - RDMPDT [3.9.1]
C.3.9.3 .. YPOSUP(T/F), ZPOSUP(T/F), LENAX, LENAY, LENAZ; (0) - ROMPDT [3.9.1]
C.3.9.4 .. IHAP1(1), IMAP2(HX), JHAP1(1), JHAP2(HY), KHAP1(1), KHAP2(HZ), AMIN, AHAX, NHPZOH(5):(0) - RDHPDT (3.9.1)
C....ONE OF THE 3.9.4 LINES REQUIRED FOR EACH DEPENDENT VARIABLE
C..... TO BE HAPPED
C....END OF FIRST SET OF TRANSIENT INFORMATION
C.....READ SETS OF READ3 DATA AT EACH TINCHG UNTIL THRU (LINES 3.H1.H2)
C....END OF CALCULATION LINES FOLLOW, THRU-. TRUE.
C.3.99.1 .. THRU
C.....TEMPORAL PLOT INFORMATION
C.3.99.2 .. PLOTWP, PLOTWT, PLOTWC; ALL (T/F)
C....PLOT INFORMATION; (0) - PLOTWP [3.99] OR PLOTWT [3.99] OR PLOTWC [3.99]
C.4.1 .. IWEL, ROPLTP(T/F)
C.4.2 .. IDLAB
C.4.3 .. NTHPTO, NTHPTC, PWHIN, PWHAX, PSNIN, PSNAX, TWHIN, TWHAX, TSHIN, TSHAX, CHIN, CHAX; (0) - RDPLTP [4.1]
C.4.4 .. TO_POW_POS_TOW_TOS_COM
C .... USE AS MANY 4.4 LINES AS NECESSARY
C.4.5 .. END WITH -1. /
C....READ DATA FOR ADDITIONAL WELLS. 4.1-4.5 LINES
C.4.6 .. END WITH 0 /
C.....END OF DATA FILE
```

A set of record pairs is needed where the first is:

I.1 I1, I2, J1, J2, K1, K2;

where Ii, Jj, Kk are inclusive node-index ranges in the i,j, and k directions, and the second is:

I.2 VAR1, IMOD1, [VAR2, IMOD2, VAR3, IMOD3];

where VARi is the value of the ith variable;

I.3 0/.

IMODi is the modification code for the ith variable;

1 means insert the data into the variable location;

2 means multiply the existing value of the variable by VARi;

3 means add VARi to the existing value of the variable;

4 means read in values of VARi on a node-by-node basis in the order of increasing i, then j, then k. In this case, the value of VARi in record I.2 is not used and needs to be given a value of zero. Note that this type of input is only suitable for zones of cells with a continuous sequence of nodes in the i-direction. Zones of cells whose i-direction index is a discontinuous sequence are more easily treated by using one of the first three types of modification code.

The brackets indicate that some input lists need three values of VAR and IMOD, while others need only one. As many lines of type I.1 and I.2 are used as necessary, then the following line is entered to terminate the input:

The generic dimensions for the various parameters are indicated, and the units must be selected from table 5.2. unless the conversion factors in the HST3D program are modified. The units for output will match the units for input. The few exceptions, where the units are not combinations of the fundamental units, are indicated. The conversion factors from inch-pound units to metric units are contained in the following subprograms: READ1,

ETOM1, ETOM2. They are specified in PARAMETER statements and the variable names are in the form, CNVxxx. The conversion factors from metric units to inch-pound units are defined in the INIT1 subprogram, and the variable names are in the form CNVxxI. The unit labels are contained in variables named UNITxx and are defined in the INIT1 subprogram also. By changing the appropriate conversion factors, and unit labels the user can employ the most convenient units for a particular simulation series.

The notation (T/F) indicates a logical variable. All index ranges are inclusive unless stated otherwise. All pressure values are relative to atmospheric unless stated otherwise.

Quantity	International	
(generic	System	Inch-pound
units)	units	units
Mass (M)	kilogram (kg)	pound (1b)
Length (L)	meter (m)	foot (ft)
Time (t)	second (s)	day (d)
Temperature (T)	degree Celsius (°C)	degree Fahrenheit (°F)
Energy (E)	Joule (J) or watt-second (W-s)	British Thermal Unit (BTU)
Fluid Volumetric Flow Rate (L <sup>3</sup> /t)	liter per second (1/s)	cubic foot per day (ft <sup>3</sup> /d)
Density (M/L <sup>3</sup> )	kilogram per cubic meter (kg/m <sup>3</sup> )	pound per cubic foot (lb/ft <sup>3</sup> ) <sup>1</sup>
Velocity (L/t)	meter per second (m/s)	foot per day (ft/d)
Pressure (F/L <sup>2</sup> )	Pascal (Pa)	pound per square inch (psi)
Viscosity (M/Lt)	kilogram per meter-second (kg/m-s)	centipoise (cP) <sup>2</sup>
Diffusivity $(L^2/t)$	square meter per second (m <sup>2</sup> /s)	square foot per day (ft <sup>2</sup> /d)
Permeability (L <sup>2</sup> )	square meter (m <sup>2</sup> )	square foot (ft <sup>2</sup> )
Thermal Conductivity	Watt per meter-degree	British Thermal Unit
(E/L-t-T) or (F/t-T)	Celsius (W/m-°C)	per foot-hour-degree Fahrenheit (BTU/ft-h-°F)
Specific Heat	Joule per kilogram-	British Thermal Unit
Capacity (FL/M-T or E/M-T)	degree Celsius	per pound-degree
	(J/kg-°C)	Fahrenheit (BTU/1b-°F)

Table 5.2.--International System and inch-pound units used in the heat- and solute-transport simulator

Quantity (generic units)	International system units	Inch-pound units
Heat-Transfer Coefficient (E/t-L <sup>2</sup> -T)	Watt per square meter- degree Celsius (W/m <sup>2</sup> -°C)	British Thermal Unit per hour-square foot- degree Fahrenheit (BTU/h-ft <sup>2</sup> -°F)
Volumetric flux (L <sup>3</sup> /L <sup>2</sup> -t)	cubic meter per square meter-second (m <sup>3</sup> /m <sup>2</sup> -s)	cubic foot per square foot-day (ft <sup>3</sup> /ft <sup>2</sup> -d)
Heat flux (E/L <sup>2</sup> -t)	Watt per square meter (W/m <sup>2</sup> )	British Thermal Unit per square foot-hour (BTU/ft <sup>2</sup> -h)
Mass flux (M/L <sup>2</sup> -t)	kilogram per square meter-second (kg/m <sup>2</sup> -s)	pound per square foot- .day (lb/ft <sup>2</sup> -d)

Table 5.2.--International System and inch-pound units used in the heat- and solute-transport simulator--Continued

<sup>1</sup>A weight density rather than a mass density. <sup>2</sup>Not Inch-Pound but common usage.

## 5.2.2. Input Record Descriptions

The following order generally has been observed for data input: (1) Fundamental and dimensioning information, (2) spatial geometry and mesh information, (3) fluid properties, (4) porous medium properties, (5) source information, (6) boundary condition information, (7) initial condition information, (8) calculation parameters, (9) output specifications. Items 5, 6, 8 and 9 have transient data associated with them, and data are input in the item-order given. The static data are read only once while the transient data are read at each time a change in the data is to occur. Only the data that are being changed need to be entered, because any unmodified data will remain the same over the next time interval of simulation. Each input record number identifies a particular record in the input-data form listed in table 5.1. Dimensioning data for space allocation - READ1

1.1 Title line 1.

A character string of up to 80 characters. It is wise to start in column two in case the string begins with the letter 'c'.

1.2 Title line 2.

A character string of up to 80 characters. It is wise to start in column two in case the string begins with the letter 'c'.

1.3 RESTRT (T/F), TIMRST.

RESTRT - True if this run is to be a restart of a previous simulation.

TIMRST - Time within the range of a previous simulation from which this restart is to continue (t). A search is made of the restartrecord file for data pertaining to the time value specified or within 0.1 percent of that time. Enter zero if this is not a restart of a previous simulation.

If this is a restart run, proceed to the READ3 group for input of transient data.

1.4 HEAT, SOLUTE, EEUNIT, CYLIND, SCALMF; all (T/F).

HEAT - True if heat transport is to be simulated.
SOLUTE - True if solute transport is to be simulated.
EEUNIT - True if the input data are in inch-pound units; otherwise,
metric units are assumed. The conversion factors are set for the
inch-pound and metric units given in table 5.1 and appear after

- the table of contents also. The program uses metric units internally.
- CYLIND True if cylindrical coordinates with no angular dependence are to be used; otherwise, cartesian, x,y,z, coordinates are used.

SCALMF - True if a scaled-mass fraction determined by equation 2.2.1.2 is to be used for data input and output; otherwise, the unscaled mass fraction will be used.

1.5 NX, NY, NZ, NHCN.

- NX The number of nodes in the x-direction for cartesian coordinates, or r-direction for cylindrical coordinates.
- NY The number of nodes in the y-direction. Unused for cylindrical coordinates, but a space in the input record must be included.
   NZ Number of nodes in the z-direction.
- NHCN Number of nodes for the heat-conduction boundary condition. This may be included only in a heat-transport simulation. These nodes are used for the finite-difference solution of equations 2.5.5.3a-d and 2.5.5.4a-d.
- 1.6 NPTCBC, NFBC, NAIFC, NLBC, NHCBC, NWEL.
  - NPTCBC Number of cells (nodes) with a specified pressure, temperature, and(or) mass-fraction boundary condition.
    - NFBC Number of cells with a specified-flux boundary condition; flow, heat, and(or) solute.
    - NAIFC Number of cell faces with an aquifer-influence-boundary condition.
    - NLBC Number of cell faces with a leakage-boundary condition.

NHCBC - Number of cell faces with a heat-conduction boundary condition.

NWEL - Number of wells in the simulation region.

These values must be the <u>exact counts</u> for the simulation. A cell with more than one face on the region boundary may be counted more than once for the above counts. A specified-value boundary-condition cell may <u>not</u> have any other type of boundary condition. 1.7 NPMZ.

NPMZ - Number of porous medium zones in the region. Porous medium hydraulic, thermal-, and solute-transport properties vary by zone.

1.8 SLMETH, LCROSD (T/F).

SLMETH - Solution method (integer):

Enter 1 to select the direct, D4, matrix-equation solver; or Enter 2 to select the iterative, two-line, successive-overrelaxation, matrix-equation solver.

LCROSD - True if the off-diagonal cross-dispersive terms in the heat and solute equations are to be lumped into the diagonal terms to give amplified coefficients. Otherwise, the coefficients in the dispersion tensor are computed explicitly in time. Lumping the coefficients reduces storage requirements and the number of iterative cycles of the flow, heat, and solute equations, but it is an empirical simplification.

1.9 IBC by i,j,k range for excluded cells.

IBC - Index of cells excluded from the simulation region. For space allocation, the location of the excluded cells must be known. Excluded cells are denoted by IBC = -1.

1.10 RDECHO (T/F).

RDECHO - True if a file is to be written that echos the input data as they are read that is used for locating data-input errors.

Static data - READ2.

The following data are invariant throughout the simulation.

2.1 PRTRE (T/F).

PRTRE - True if a read-echo printout of data input by i,j,k range is desired.

The following 12 records describe the gridding of the simulation region.

2.2A.1 UNIGRX, UNIGRY, UNIGRZ; All (T/F); optional, used for cartesiancoordinate systems.

UNIGRi - True if the grid in the ith direction is uniform.

2.2A.2A X(1),X(NX); Optional, used if uniform grid in the x-direction.

X(1) - Location of the first node point in the x-direction (L). X(NX) - Location of the last node point in the x-direction (L).

2.2A.2B X(I), I=1 to NX; Optional, used for nonuniform grid in the x-direction.

2.2A.3A Y(1), Y(NY); Optional, used if uniform grid in the y-direction.

Y(1) - Location of the first node point in the y-direction (L). Y(NY) - Location of the last node point in the y-direction (L).

2.2A.3B Y(J) J=1 to NY; Optional, used for nonuniform grid in the y-direction.

2.2A.4A Z(1), Z(NZ); Optional, used for uniform grid in the z-direction.

Z(1) - Location of the first node point in the z-direction (L). Z(NZ) - Location of the last node point in the z-direction (L).

2.2A.4B Z(K), K=1 to NZ; Optional, used for nonuniform grid in the z-direction.

- 2.2B.1A R(1), R(NR), ARGRID (T/F); Optional, used for cylindrical-coordinate systems.
  - R(1) Interior radius of cylindrical-coordinate system (L). Required by cylindrical-coordinate system.
  - R(NR) Exterior radius of cylindrical-coordinate system (L). Required if a cylindrical-coordinate system.
  - ARGRID True if automatic gridding or node location in the r-direction is desired. A logarithmic spacing in R will be used according to equation 3.1.2.1.
- 2.2B.1B R(I), I = 1 to NX; Optional, used for user-specified radial gridding for a cylindrical-coordinate system.
  - R(I) Array of node locations along the r-axis (L). Required by cylindrical-coordinate system if automatic gridding is not selected.

2.2B.2 UNIGRZ (T/F).

- 2.2B.3A Z(1), Z(NZ); Optional, used if uniform grid in the z-direction for cylindrical coordinates.
- 2.2B.3B Z(K), K = 1 to NZ; Optional, used for nonuniform grid in the z-direction for cylindrical coordinates.

Z(K) - Array of node locations along the z-axis (L).

The following two records describe a tilted-coordinate system.

- 2.3.1 TILT (T/F); Optional, required only if a cartesian-coordinate system is used.
  - TILT True if a tilted-coordinate system is desired with the z-axis not in the vertical upward direction.

No tilt may be specified if a free-surface boundary condition is to be employed (Record 2.20).

- 2.3.2 THETXZ, THETYZ, THETZZ; Optional, required only if a tiltedcoordinate system is being used.
  - THETXZ Angle that the x-axis makes with the vertical upward direction (DEG).
  - THETYZ Angle that the y-axis makes with the vertical upward direction (DEG).

THETZZ - Angle that the z-axis makes with the vertical upward direction (DEG).

Fluid properties

2.4.1 BP.

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BP - Compressibility of the fluid  $(F/L^2)^{-1}$ .

Fluid density data

2.4.2 PRDEN, TRDEN, WO, DENFO.

. PRDEN - Reference pressure for density (relative to atmospheric) (eq. 2.2.1.1b) (F/L<sup>2</sup>).

TRDEN - Reference temperature for density (eq. 2.2.1.1b) (T).

WO - Reference mass fraction for density (eq. 2.2.1.1b) and minimum-mass fraction for scaling (eq. 2.2.1.2) and  $\rho_0\beta_W$  term (eq. 2.2.1.1c) (-). Needs to be zero if solute decay takes place. Should be zero if solute transport is not being simulated.

DENFO - Fluid density at the minimum solute mass fraction (eq. 2.2.1.1c or eq. 2.2.1.3b) (M/L<sup>3</sup>). (kg/m<sup>3</sup>) or (lb/ft<sup>3</sup>).

- 2.4.3 W1, DENF1; Optional, required only if a solute transport is being simulated.
  - W1 Maximum-mass fraction for scaling (eq. 2.2.1.2) and  $\rho_{0}\beta_{W}$  term (eq. 2.2.1.1c) (-). Should be equal to or greater than the maximum mass-fraction value specified by any boundary condition or initial condition.
  - DENF1 Fluid density at the maximum solute mass fraction (eq. 2.2.1.1c or eq. 2.2.1.3b) (M/L<sup>3</sup>) (kg/m<sup>3</sup>) or (lb/ft<sup>3</sup>).

If solute transport is being simulated and a scaled-mass fraction has been selected for input and output (SCALMF in record 1.4), WO and WI are used to perform the scaling according to equation 2.2.1.2 and slope calculation of equation 2.2.1.3b.

Fluid-viscosity data

- 2.5.1 NOTVO, (TVFO(I), VISTFO(I), I = 1 to NOTVO); Optional, required if only flow; or flow and heat transport; or flow, heat and solute transport are being simulated. Not used if only flow and solute transport are being simulated.
  - NOTVO Number of viscosity versus temperature points for fluid at minimum or reference solute-mass fraction (minimum of one point).
     TVFO - Array of temperature points (T).
     VISTFO - Array of viscosity points at minimum solute-mass fraction, WO,
- 2.5.2 NOTV1 (TVF1(I), VISTF1(I) I = 1 to NOTV1); Optional, required only if heat and solute transport are being simulated.

(M/Lt) (kg/m-s) or (cP).

 NOTV1 - Number of viscosity versus temperature points for fluid at maximum solute mass fraction (minimum of one point).
 TVF1 - Array of temperature points (T).
 VISTF1 - Array of viscosity points at maximum solute-mass fraction, W1, (M/Lt), (kg/m-s) or (cP). 2.5.3 NOCV, TRVIS, (CVIS(I), VISCTR(I), I = 1 to NOCV); Optional, required only if solute transport is being simulated.

NOCV - Number of viscosity versus mass-fraction points (minimum of two points).

TRVIS - Reference temperature for viscosity versus mass-fraction data (T).
CVIS - Array of mass-fraction (or scaled-mass-frac: ion) points (-).
VISCTR - Array of viscosity points for fluid at reference temperature
(M/Lt), (kg/m-s) or (cP).

If only solute transport is being simulated, only record number 2.5.3 is needed from this group.

Reference condition information

2.6.1 PAATM.

PAATM - Atmospheric absolute-pressure value used to relate gage pressure to absolute pressure  $(F/L^2)$ .

If zero is entered, standard atmospheric pressure of 1.01325×10<sup>5</sup> Pa is used.

2.6.2 POH, TOH.

- POH Reference pressure (relative to atmospheric) for enthalpy variations,  $P_{OH}$ , (F/L<sup>2</sup>). This value should be within the range of pressure to be simulated.
- TOH Reference temperature for enthalpy variations or the constant temperature for isothermal simulations, T<sub>oH</sub> (T). This value should be within the range of temperature to be simulated.

These values are used in equation 2.2.3.1c.

### Fluid thermal properties

- 2.7 CPF, KTHF, BT; Optional, required only if heat transport is being simulated.
  - CPF Fluid heat capacity at constant pressure (FL/M-T). An average value for the range of solute concentration and temperature to be simulated should be used (eq. 2.2.3.1b and eq. 2.2.3.1c).
  - KHTF Fluid thermal conductivity (F/t-T). An average value for the range of solute concentration and temperature to be simulated should be used.
  - BT Fluid coefficient of thermal expansion (T<sup>-1</sup>). An average value for the range of solute concentration and temperature to be simulated should be used.

Solute information

- 2.8.1 DM, DECLAM; Optional, required only if solute transport is being simulated.
  - DM Effective molecular diffusivity for the solute in the porous media  $(L^2/t)$ . DECLAM - Solute-decay-rate constant  $(t^{-1})$ .

The following two records describe the zonation of the simulation region.

2.9.1 IPMZ, I1Z(IPMZ) I2Z(IPMZ), J1Z(IPMZ), J2Z(IPMZ), K1Z(IPMZ), K2Z(IPMZ).

IMPZ - Porous-medium zone number.

Records of this form define the zones within the simulation region and assign zone numbers. These zones are used to assign values to the porousmedium properties. The ranges of the indices in the I, J, and K directions define the rectangular prism for a given zone. The zones must be convex and non-overlapping as explained in section 3.1 and the entire simulation region must be covered by the set of zones. No zones may be defined that include elements outside the simulation region. For cylindrical coordinates, J1Z and J2Z must equal 1. The subscript IPMZ identifies data that are input by zone number in subsequent records. The number of 2.9.1 records must equal NPMZ.

2.9.2 End the input with 0 / .

Porous-media properties

2.10.1 KXX(IPMZ), KYY(IPMZ), KZZ(IPMZ); IPMZ = 1 to NPMZ.

KXX - Permeability in the x-direction or r-direction for zone IPMZ  $(L^2)$ .

KYY - Permeability in the y-direction for zone IPMZ (L<sup>2</sup>). Not used for cylindrical coordinates, but a zero or blank space must be indicated in the input record.

KZZ - Permeability in the z-direction for zone IPMZ  $(L^2)$ .

2.10.2 POROS(IPMZ), IPMZ = 1 to NPMZ.

POROS - Porosity of the medium in zone IPMZ (-).

2.10.3 ABPM(IPMZ), IPMZ = 1 to NPMZ.

ABPM - Porous-medium bulk vertical compressibility in zone IPMZ (F/L<sup>2</sup>)<sup>-1</sup>. This compressibility is determined on a fixed mass of porous medium undergoing vertical compression.

Porous-media thermal properties

- 2.11.1 RCPPM(IPMZ), IPMZ = 1 to NPMZ; Optional, required only if heat transport is being simulated.
  - RCPPM Heat capacity of the porous-medium solid phase per unit volume for zone IPMZ  $(F/L^2-T)$ .

- 2.11.2 KTXPM(IPMZ), KTYPM(IPMZ), KTZPM(IPMZ), IPMZ = 1 to NPMZ; Optional, required only if heat transport is being simulated.
  - KTXPM Thermal conductivity of the porous medium in the x-direction for zone IPMZ (F/t-T).
  - KTYPM Thermal conductivity of the porous medium in the y-direction for zone IPMZ (F/t-T). Not used if cylindrical coordinates, but a zero or a blank space in the input file must be denoted.
  - -KTZPM Thermal conductivity of the porous medium in the z-direction for zone IPMZ (F/t-T).

Solute and thermal dispersion information

2.12 ALPHL(IPMZ), ALPHT(IPMZ), IPMZ = 1 to NPMZ; Optional, required only if heat or solute transport is being simulated.

ALPHL - Longitudinal dispersivity for zone IPMZ (L). ALPHT - Transverse dispersivity for zone IPMZ (L).

Solute sorption information

- 2.13 DBKD(IPMZ) IPMZ = 1 to NPMZ; Optional, required only if solute transport is being simulated.
  - DBKD Dimensionless linear-equilibrium-distribution coefficient. This is the porous-medium bulk density times the distribution coefficient (-).

Well-model information

2.14.1 RDWDEF (T/F); Optional, required only if there are wells in the simulation region.

RDWDEF - True if well definition data are to be read.

- 2.14.2 IMPQW (T/F); Optional, required only if there are wells in the region and cartesian coordinates are being used.
  - IMPQW True if semi-implicit well-flow calculations are to be made. Otherwise, well-flow rates are calculated explicitly at the beginning of the time step.

Record lines 2.14.3-6 are repeated as often as necessary for each well in the simulation. Each well must be defined at this point, whether or not it is active at the start of the simulation.

2.14.3 IWEL, IW, JW, LCBOTW, LCTOPW, WBOD, WQMETH; Optional, required only if there are wells in the simulation region and well-definition data are to be read.

IWEL - Well number.

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- IW Cell number in x-direction of well location.
- JW Cell number in y-direction of well location. Not used in cylindrical coordinates, but a zero or a blank entry must be included.

LCBOTW - Cell number in z-direction of lowermost completion layer for the well.

LCTOPW - Cell number in z-direction of uppermost completion layer for the well. This is the location at which the dependentvariable data are taken if this is an observation well.

WBOD - Well bore outside diameter. This is the drilled diameter or diameter of the screen or perforated casing (L).

WQMETH - Index for well-flow calculation method (integer).

10 - Specified well-flow rate with allocation by mobility and pressure difference.

11 - Specified well-flow rate with allocation by mobility.

20 - Specified pressure at well datum with allocation by mobility and pressure difference.

- 30 Specified well-flow rate with a limiting pressure at well datum. Flow-rate allocation by mobility and pressure difference.
- 40 Specified surface pressure with allocation by mobility and pressure difference. Well-riser calculations will be performed.
- 50 Specified surface-flow rate with limiting surface pressure. Allocation by mobility and pressure difference. Well-riser calculations will be performed.
- 0 Observation well or abandoned well.
- 2.14.4 WCF(L), L = 1 to NZ-1; Optional, required only if there are wells in the simulation region and well-definition data are to be read.
  - WCF Well-completion-factor.array on an element-by-element basis (-). Element L goes from the node at z-level L to the node at z-level L+1. An element completion factor of one means the geometric-mean of the horizontal permeability for that element will be used in the well index. An element completion factor of zero means the well is cased off from the aquifer in that element. A reduced permeability around the well bore can be approximately represented by specifying a completion factor less than one. Note that this is not the same as a well skin factor, which is not included in the present version of the program. If WCF is zero for the element between LCBOTW and LCBOTW+1 or for the element between LCTOPW-1 and LCTOPW, then LCBOTW or LCTOPW should be adjusted as necessary to range from the bottom to the top completion layer that communicate with the aquifer. The well-completion factor also can be used to compute an approximate effective permeability for a well that is completed in a cell that contains multiple zones of different permeability.
- 2.14.5 WRISL, WRID, WRRUF, WRANGL; Optional, required only if welldefinition data are to be read and well-riser calculations are to be made (WQMETH is 40 or 50).

WRISL - Well riser-pipe length (L).

WRID - Well riser-pipe inside diameter (L).

WRRUF - Well riser-pipe roughness factor (L) (see table 2.1). WRANGL - Well riser-pipe angle with vertical direction (DEG).

- 2.14.6 HTCWR, DTHAWR, KTHAWR, KTHWR, TABWR, TATWR; Optional, required only for heat-transport simulations, and if well-riser calculations are to be done.
  - HTCWR Heat-transfer coefficient from the fluid to the well-riser pipe  $(E/tL^2T)$ .
  - DTHAWR Thermal diffusivity of the medium adjacent to the well-riser pipe  $(L^2/t)$ .
  - KTHAWR Thermal conductivity of the medium adjacent to the well-riser pipe (F/t-T).
  - KTHWR Thermal conductivity of the well-riser pipe (F/t-T).

TABWR - Ambient temperature at the bottom of the well-riser pipe (T).

TATWR - Ambient temperature at the top of the well-riser pipe (T).

2.14.7 End this data set with 0/.

- 2.14.8 MXITQW, TOLDPW, TOLFPW, TOLQW, DAMWRC, DZMIN, EPSWR; Optional, required only if wells are in the region, and if well-riser calculations are to be done.
  - MXITQW Maximum number of iterations allowed for the well-flow rate allocation calculation. Default of 20.
  - TOLDPW Tolerance on the change in well-riser pressure for the wellriser iterative calculation ( $F/L^2$ ). Default of  $6 \times 10^{-3}$  Pa. This is the primary convergence test.
  - TOLFPW Tolerance on the fractional change in well-riser pressure for the well-riser iterative calculation. Default of 0.001. This is the secondary convergence test.

- TOLQW Tolerance on the fractional change in well flow rate because of temperature and mass-fraction changes for the source term in the flow equation. Default of 0.001. This is the tertiary convergence test.
- DAMWRC Damping factor for well-pressure adjustment during the iterations for allocating flow rates. Default of 2.
- DZMIN Minimum value of step length along the well riser (L). Default of 1 percent of riser length.
- EPSWR Fractional tolerance for the integration of the pressure and temperature equations along the well riser. Default of 0.001.

Boundary-condition information Specified value

- 2.15 IBC by i,j,k range; Optional, required only if specified value boundary-condition cells are present.
  - IBC Index of boundary-condition type. This is in the form  $n_1n_2n_3$ where  $n_1$  refers to pressure;  $n_2$  refers to temperature; and  $n_3$  refers to mass fraction. The value for  $n_1$  is set to 1 to indicate that a specified value boundary condition for variable i exists at that cell.

Remember that a specified value for cell removes that cell from the calculation. Therefore, no other boundary conditions at that cell can be specified for the equation concerned.

#### Specified flux

- 2.16 IBC by i,j,k range; Optional, required only if specified flux boundarycondition cells are present.
  - IBC Index of boundary-condition type. This is in the form  $n_100n_4n_5n_6$ where  $n_1=1,2,3$ , meaning that the flux is through the x,y, or z boundary face respectively. Values for  $n_4$ ,  $n_5$ , and  $n_6$  are set to

2 to indicate that a specified flux boundary condition, for a chosen equation, is required at that cell, where  $n_4$  refers to the flow equation,  $n_5$  refers to the heat-transport equation, and  $n_6$  refers to the solute-transport equation.

Aquifer leakage

- 2.17.1 IBC by i,j,k range for aquifer-leakage boundary cells; Optional, required only if leakage boundary conditions are being used.
  - IBC Index of boundary-condition type. It is of the form  $n_100300$  for aquifer leakage, where  $n_1$  indicates the direction of the normal to the leakage boundary face. Values for  $n_1$  are 1 for the x-direction; 2 for the y-direction; and 3 for the z-direction. The number 3 in the hundreds place denotes an aquifer-leakage boundary condition.
- 2.17.2 KLBC, BBLBC, ZELBC by i,j,k range; Optional, required only if leakage boundary conditions are being used.
  - KLBC Permeability of confining layer for aquifer-leakage boundary condition (L<sup>2</sup>). Appears in equations 2.5.3.1a and 3.4.3.1a.
    BBLBC Confining-layer thickness for leakage boundary condition (L).
    ZELBC Elevation of the far side of the confining layer away from the simulation region (L).

For leakage across lateral boundaries of the simulation region, ZELBC is automatically set equal to the elevation of the corresponding boundary node.

River leakage

2.17.3 I1,I2,J1,J2, KRBC, BBRBC, ZERBC; Optional, required only if the river-leakage boundary condition is being used.

- I1,I2,J1,J2 Node or cell number ranges in the x and y directions for a river-leakage boundary-condition segment. River segments are lines in the x-, y-, or diagonal direction.
  - KRBC Permeability multiplied by effective-riverbed-area factor for river-leakage boundary-condition (L<sup>2</sup>). The effectiveriverbed-area factor is the ratio of the riverbed area to the boundary-face area for a given cell.
  - BBRBC Thickness of the confining layer that forms the riverbed (L). ZERBC - Elevation of the top of the confining layer that forms the riverbed defined in figure 2.4 (L).

Use as many 2.17.3 records as necessary to describe the river.

2.17.4 End this data set (records 2.17.3) with 0/.

Aquifer-influence functions

- 2.18.1 IBC by i,j,k range for aquifer-influence-function boundary cells; Optional required only if aquifer-influence-function boundary conditions are being used.
  - IBC Index of boundary-condition type. It is in the form  $n_100400$ for aquifer-influence functions, where  $n_1$  indicates the direction of the normal to the influence-function-boundary face. Values for  $n_1$  are 1 for the x-direction; 2 for the y-direction; and 3 for the z-direction. The number 4 in the hundreds place denotes an gauifer-influence-function boundary condition.
- 2.18.2 UVAIFC by i,j,k range; Optional, required only if aquifer-influence functions are used.
  - UVAIFC Temporary storage for input of user-specified factors for aquifer-influence-function spatial allocation.

These factors are defined on a zonal basis, so the factor for all boundary zones which include a common cell for which an aquifer-influencefunction boundary condition applies must be the same. If default weighting of the aquifer-influence functions in proportion to their boundary-cell facial area is desired, no values for UVAIBC need to be entered (except the closing 0 /).

1.1.1

2.18.3 IAIF; Optional, required only if aquifer-influence functions are being used.

IAIF - Index of aquifer-influence function.

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1 - Pot aquifer for outer-aquifer region.

- 2 Transient-aquifer-influence function with calculation using the Carter-Tracy approximation.
- 2.18.4 KOAR, ABOAR, VISOAR, POROAR, BOAR, RIOAR, ANGOAR; Optional, required only if transient-aquifer-influence functions are being used.

KOAR - Permeability for the outer-aquifer region  $(L^2)$ .

- ABOAR Porous-medium bulk vertical compressibility for the outeraquifer region  $(F/L^2)^{-1}$ .
- VISOAR Viscosity of the fluid in the outer-aquifer region (M/Lt), (kg/m-s) or (cP).
- POROAR Porosity for the outer-aquifer region (-).
- BOAR Total thickness of the outer-squifer region (L).
- RIOAR Radius of the equivalent cylinder that contains the inneraquifer region (L). Usually determined by equation 3.4.4.2.1.

ANGOAR - Angle of influence of the outer-aquifer region (DEG.).

This is the angle subtended by the part of the equivalent cylindrical boundary that is subject to flux determined by the aquifer-influence function.

The following three records describe the gridding and parameters for heatconduction boundary conditions.

- 2.19.1 IBC by i,j,k range for heat-conduction boundary-condition nodes; Optional, required only if a heat simulation, and if heat-conduction boundary condition cells are to be used.
  - IBC Index of boundary-condition type, denoted by a number in the form  $n_100040$  where  $n_1$  indicates the direction of the outward normal to the heat-conduction boundary-condition cell face. The values for  $n_1$  are: 1 for the x-direction; 2 for the y-direction; and 3 for the z-direction. The signs of the normal is disregarded. The 4 indicates that this cell has a heat-conduction boundary condition on one of its faces. Only one face of a given cell can be assigned a heat-conduction boundary condition.
- 2.19.2 ZHCBC(K) K = 1 to NHCN; Optional, required if heat simulation is being done, and there are heat-conduction boundaries (NHCBC > 0).
  - ZHCBC Array of node distances along the outward pointing normal from a heat-conduction boundary surface (L). The first value must be zero. All heat-conduction boundary-condition cells use the same nodal distribution.
- 2.19.3 UDTHHC by i,j,k range for heat-conduction boundary-condition cells;
  Optional, required only if a heat simulation, and if there are heatconduction boundary-condition cells.
  - UDTHHC Temporary storage for input of thermal diffusivity of the heat-conducting medium outside the simulation region as a function of boundary-cell location  $(L^2/t)$ .
- 2.19.4 UKHCBC by i,j,k range for heat-conduction boundary-condition cells; Optional, required only if a heat simulation, and if there are heatconduction boundary-condition cells.
  - UKHCBC Temporary storage for input of thermal conductivity of the heat-conducting medium outside the simulation region as a function of boundary-cell location (E/L-t-T).

Free-surface boundary condition

2.20 FRESUR (T/F), PRTCCM (T/F).

- FRESUR True if the region is unconfined, so that a free-surface boundary exists.
- PRTCCM True if a message is to be printed when a free surface rises above the top of a cell or falls below the bottom of a cell, or if a cell below the uppermost layer becomes unsaturated.

Initial conditions

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2.21.1 ICHYDP (T/F), ICT (T/F), ICC (T/F).

- ICHYDP True if initial condition of hydrostatic pressure distribution is to be specified.
- ICT True if an initial-condition temperature distribution is to be specified.

ICC - True if an initial-condition mass-fraction distribution is to be specified.

2.21.2 ICHWT (T/F); Optional, required only if a free-surface boundary exists.

ICHWI - True if an initial-condition water-table-elevation distribution is to be input.

2.21.3A ZPINIT, PINIT; Optional, required only if an initial-condition hydrostatic-pressure distribution is being specified.

ZPINIT - Elevation of the initial-condition pressure (L). PINIT - Pressure for hydrostatic, initial-condition distribution  $(F/L^2)$ .

2.21.3B P by i,j,k range; Optional, required only if a non-hydrostatic pressure distribution is being specified as an initial condition.

P - Pressure distribution for the initial condition  $(F/L^2)$ .

- 2.21.3C HWT by i,j,k range; Optional, required only if desired in conjunction with a free-surface boundary condition.
  - HWT Water-table-elevation distribution for the initial condition (L). Specified for the upper layer of cells only.
- 2.21.4A NZTPRO (ZT(I), TVD(I); I = 1 to NZTPRO); Optional, required only if a heat simulation is being done.
  - NZTPRO Number of points in the temperature-versus-depth profile for initial-condition temperature distribution. Limit of 10.
  - 2T Array of locations along the z-axis for initial-temperature distribution (L). These locations must span the entire z-axis range of the region.

TVD - Array of initial temperatures along the z-axis (T).

2.21.4B T by i,j,k range; Optional, required only if ICT is true.

T - Temperature distribution for the initial condition (T).

- 2.21.5 NZTPHC, ZTHC(I), TVZHC(I), I = 1 to NZTPHC; Optional, required only if a heat-transport simulation is being done, and if there are heatconduction boundary conditions.
  - NZTPHC Number of points in the outward normal direction to the heatconduction boundary-condition surfaces for initial-conditiontemperature profile. Limit of 5.
  - ZTHC Array of node locations in the outward normal direction for initial-condition-temperature profile for heat-conduction boundary-condition cell faces (L). The first value must be zero, and these nodes must span the mesh defined by ZHCBC in record 2.19.2.
  - TVZHC Array for the initial-condition temperature-profile values for heat-conduction boundary-condition cell faces (T).

The same initial-condition profile is used for each heat-conduction boundary condition cell.

2.21.6 C by i, j, k range; Optional, required only if ICC is true.

C - Mass-fraction (or scaled mass fraction) distribution for the initial condition (-).

Calculation information

2.22.1 FDSMTH, FDTMTH.

FDSMTH - Factor for spatial-discretization method.

0.5 - centered-in-space differencing used for advective terms.

0 - upstream differencing in space used for advective terms.

FDTMTH - Factor for temporal-discretization method.

0.5 - centered-in-time or Crank-Nicholson differencing used.

1. - backward-in-time or fully-implicit differencing used.

2.22.2 TOLDEN, MAXITN.

- TOLDEN Tolerance in fractional change in density for convergence over a solution cycle of flow, heat, and solute equations at a given time plane. Default set at 0.001.
- MAXITN Maximum number of iterations allowed for a cycle of pressure, temperature, and mass-fraction solutions allowed at a given time plane. Default set at 5.

2.22.3 NTSOPT, EPSSOR, EPSOMG, MAXIT1, MAXIT2; Optional, required only if the two-line, successive-overrelaxation method for solving the system matrix equations is selected.

- NTSOPT Number of time steps between recalculations of the optimumoverrelaxation parameter. Default set at 5.
- EPSSOR Tolerance for the two-line, successive-overrelaxation iterative solution of the matrix equations at each time plane. Default set at  $1 \times 10^{-5}$ . The maximum fractional change in any of the values of the dependent variable must be less than or equal to this tolerance times  $(2-w_{opt})$ .
- EPSOMG Tolerance on the fractional change in the overrelaxation parameter during the iterative calculation to determine the optimum value. Default set to 0.2.
- MAXIT1 Maximum number of iterations allowed for the calculation of the optimum overrelaxation parameter. Default set at 50.
- MAXIT2 Maximum number of iterations allowed for the solution of the matrix equations. Default set at 100.

Output of static data

2.23.1 PRTPMP (T/F), PRTFP (T/F), PRTIC (T/F), PRTBC (T/F), PRTSLM (T/F), PRTWEL (T/F).

PRTPMP - True if a printout of porous-media properties is desired.
PRTFP - True if a printout of fluid properties is desired.
PRTIC - True if a printout of initial conditions is desired.
PRTBC - True if a printout of static boundary-condition information is desired.
PRTSLM - True if a printout of solution-method information is desired.
PRTWEL - True if a printout of static-well bore information is desired.

2.23.2 IPRPTC, PRTDV (T/F); Optional, required only if initial-condition printouts of the dependent variables are desired.

IPRPTC - Index of printout for initial-condition information. It is of the form  $n_1n_2n_3$ , where  $n_i$  is set to 1 for printout of the ith variable, otherwise  $n_i$  is set to 0. The variables are  $n_1$ for pressure;  $n_2$  for temperature; and  $n_3$  for mass fraction. In addition,  $n_1$  is set to 2 for both pressure and potentiometric head to be printed for isothermal cases;  $n_2$  is set to 2 for both temperature and fluid enthalpy to be printed. PRTDV - True if a printout of the density and viscosity arrays is desired.

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2.23.3 ORENPR; Optional, required only for a cartesian-coordinate system.

ORENPR - Index for orientation of the array printouts (integer);

- 12 Means x-y printouts for each plane along the z-axis, areal layers.
- 13 Means x-z (or r-z) printouts for each plane along the y-axis, vertical slices.

A negative value means the y or z-axis is positive down the page.

2.23.4 PLTZON (T/F); Optional, required only if printout of porous-media properties has been requested.

PLTZON - True if a line-printer plot of the porous-media property zones is desired.

2.23.5 OCPLOT (T/F).

OCPLOT - True if plots of observed and calculated values of the dependent variables are to be plotted versus time at the end of the simulation.

Transient data - READ3

Groups of transient data are read by subroutine READ3; one at the beginning and others during the simulation, as necessary, whenever sources, boundary conditions, calculation parameters, or output options are to be changed. Only the parameters that are to be changed need to be input. The remaining parameters will keep their previous values.

3.1 THRU (T/F).

THRU - True if the simulation is through, and the closing procedures can begin. Proceed to record 3.99 if the simulation is finished.

Well information

3.2.1 RDWFLO (T/F), RDWHD (T/F); Optional, required only if there are wells in the simulation region.

RDWFLO - True if well-flow-rate data is to be read at this time. RDWHD - True if well-head data is to be read at this time.

3.2.2 IWEL, QWV, PWSUR, PWKT, TWSRKT, CWKT; Optional, required only if well-flow or well-head data are to be read at this time.

IWEL - Well number.

QWV - Volumetric flow rate for this well  $(L^3/t)$ , (2/s) or  $(ft^3/d)$ .

PWSUR - Pressure at the land surface for this well  $(F/L^2)$ .

- Used when surface conditions are specified and the well-riser calculation is to be done.
- PWKT Pressure at the well datum for this well (F/L<sup>2</sup>). Used when well-datum conditions are specified, and no well-riser calculation is to be done.
- TWSRKT Fluid temperature at the land surface or well datum for this well (T). Used when surface conditions are specified for an injection well, and used for the well-datum value, when well-datum conditions are specified.

CWKT - Mass fraction (or scaled-mass fraction) at the well datum for this well (-). Surface and well-datum concentrations are equal, so this variable also is used to specify surface conditions for an injection well.

As many records of type 3.2.2 are used as necessary to define conditions at all the wells. Data do not have to be input for any well that does not have its conditions changed at this time.

3.2.3 End this data set with 0/.

Boundary-condition information

Specified value

3.3.1 RDSPBC (T/F), RDSTBC (T/F), RDSCBC (T/F); Optional, required only if there are specified-pressure, temperature or mass-fraction boundarycondition cells.

RDSPBC - True if specified-pressure boundary-condition data are to be read at this time.

RDSTBC - True if specified-temperature boundary-condition data are to be read at this time.

RDSCBC - True if specified mass-fraction boundary-condition data are to be read at this time.

3.3.2 PNP by i,j,k range; Optional, required only if specified-pressure boundary-condition values are to be input.

PNP - Pressure at specified-pressure boundary-condition nodes  $(F/L^2)$ .

3.3.3 TSBC by i,j,k range; Optional, required only if specified-pressure boundary-condition values are to be read, and if a heat-transport simulation is being done.

- TSBC Temperature associated with a specified-pressure boundary condition node (T). If inflow occurs, this temperature will determine the heat-inflow rate.
- 3.3.4 CSBC by i,j,k range; Optional, required only if specified-pressure boundary-condition values are to be read, and if solute transport is being simulated.
  - CSBC Mass fraction (or scaled-mass fraction) associated with a specified-pressure boundary-condition node (-). If inflow occurs, this mass fraction will determine the solute-inflow rate.
- 3.3.5 TNP by i,j,k range; Optional, required only if specified-temperature boundary-condition data are to be read.
  - TNP Temperature at specified-temperature boundary-condition nodes (T).
- 3.3.6 CNP by i,j,k range; Optional, required only if specified massfraction values are to be input.
  - CNP Mass fraction (or scaled-mass fraction) for specified massfraction boundary-condition nodes.

## Specified flux

- 3.4.1 RDFLXQ (T/F), RDFLXH (T/F), RDFLXS (T/F); Optional, required only if specified-flux boundary conditions exist.
  - RDFLXQ True if specified fluid-flux values are to be read at this time.
  - RDFLXH True if specified heat-flux values are to be read at this time.

RDFLXS - True if specified solute-flux values are to be read at this time.

- 3.4.2 QFFX, QFFY, QFFZ by i,j,k range; Optional, required only if specified fluid-flux values are to be read at this time.
  - QFFX, QFFY, QFFZ Components of the fluid-flux vector for a boundary cell in the x,y, and z-coordinate directions, respectively  $(L^3/L^2t)$ .
- 3.4.3 UDENBC by i,j,k range; Optional, required only if specified fluidfluxes values are to be read at this time.
  - UDENBC Density associated with specified-fluid flux (M/L<sup>3</sup>); (kg/m<sup>3</sup>) or (1b/ft<sup>3</sup>). If inflow, this density determines the mass flux.
- 3.4.4 TFLX by i,j,k range; Optional, required only if specified fluid fluxes are to be read at this time, and if heat transport is being simulated.
  - TFLX Temperature associated with specified fluid flux (T). If inflow, this temperature determines the heat flux.
- 3.4.5 CFLX by i,j,k range; Optional, required only if specified fluid fluxes are to be read at this time, and if solute transport is being simulated.
  - CFLX Mass fraction (or scaled mass fraction) associated with specified fluid flux (-). If inflow, this mass fraction determines the solute flux.
- 3.4.6 QHFX, QHFY, QHFZ by i,j,k range; Optional, required only if specified heat-flux values are to be read at this time.

- QHFX, QHFY, QHFZ Components of the specified heat-flux vector for a boundary cell in the x,y and z-coordinate directions respectively. Heat flux should be specified only through faces where there is no fluid flux  $(E/L^2-t)$ .
- 3.4.7 QSFX, QSFY, QSFZ by i,j,k range; Optional, required only if specified solute-flux values are to be read at this time.
  - QSFX, QSFY, QSFZ Components of the specified solute flux for a boundary cell in the x,y and z coordinate directions respectively. Solute flux should be specified only through faces where there is no fluid flux (M/L<sup>2</sup>-t).

Leakage boundary conditions

- 3.5.1 RDLBC(T/F); Optional, required only if leakage boundary-condition cells are employed.
  - RDLBC True if leakage boundary-condition data are to be read this time.
- 3.5.2 PHILBC, DENLBC, VISLBC by i,j,k range; Optional, required only if leakage boundary-condition data are to be read at this time.
  - PHILBC Potential energy per unit mass of fluid (eq. 2.5.3.1.1b) on the other side of the aquitard from the simulation region (E/M).
  - DENLEC Density of the fluid on the other side of the aquitard  $(M/L^3)$ ,  $(kg/m^3)$  or  $(lb/ft^3)$ .
  - VISLBC Viscosity of the fluid on the other side of the aquitard (M/L-t); (kg/m-s) or (cP).
- 3.5.3 TLBC by i,j,k range; Optional, required only if leakage boundarycondition data are to be read at this time, and if heat transport is being simulated.

TLBC - Temperature of the fluid on the other side of the aquitard (T).

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3.5.4 CLBC by i,j,k range; Optional, required only if leakage boundarycondition data are to be read at this time, and if solute transport is being simulated.

CLEC - Solute-mass fraction (or scaled-mass fraction) on the other side of the aquitard (-).

River leakage

- 3.5.5 I1,I2,J1,J2, HRBC, DENRBC, VISRBC, TRBC, CRBC; Optional, required only if river-leakage boundary-condition data are to be read at this time.
  - I1,I2,J1,J2 Node or cell number ranges in the x and y directions for a river-leakage boundary-condition segment. They should correspond to the segments used to define the river in data record 2.22.4.

HRBC - Potentiometric head in the river (L).

- DENRBC Density of the river fluid  $(M/L^3)$ ;  $(kg/m^3)$  or  $(1b/ft^3)$ .
- VISRBC Viscosity of the river fluid (M/L-t); (kg/m-s) or (cP).

TRBC - Temperature of the river fluid (T).

CRBC - Solute-mass fraction (or scaled-mass fraction) of the river fluid (-).

As many records of type 3.5.5 are used as necessary to include all the cells at which a river-leakage boundary condition exists.

3.5.6 End this data set (record 3.5.5) with 0 /.

Aquifer influence functions

3.6.1 RDAIF (T/F); Optional, required only if aquifer-influence-function boundary-condition cells are employed.

- RDAIF True if aquifer-influence-function boundary-condition data are to be read at this time.
- 3.6.2 DENOAR by i,j,k range; Optional, required only if aquifer-influencefunction cells are employed.
  - DENOAR Density of the fluid in the outer-aquifer region  $(M/L^3)$ ;  $(kg/m^3)$  or  $(lb/ft^3)$ .
- 3.6.3 TAIF by i,j,k range; Optional, required only if aquifer-influencefunction boundary condition cells are employed, and if heat transport is being simulated.
  - TAIF Temperature of the fluid in the outer-aquifer region associated with a given aquifer-influence-function cell (T).
- 3.6.4 CAIF by i,j,k range; Optional, required only if aquifer-influencefunction cells are employed, and if solute transport is being simulated.
  - CAIF Mass fraction of solute in the outer-aquifer region associated with a given aquifer-influence-function cell (-).

Calculation information

The following data pertains to time-step control and the time when new transient data will be read.

3.7.1 RDCALC (T/F).

RDCALC - True if calculation information is to be read at this time.

3.7.2 AUTOTS (T/F); Optional, required only if calculation information is to be read at this time.

AUTOTS - True if automatic time-step adjustment is desired for the next interval of simulation time.

3.7.3A DELTIM; Optional, required only if automatic time-step calculation is not being used and calculation information is being read at this time.

DELTIM - Time-step length (t).

- 3.7.3B DPTAS, DTTAS, DCTAS, DTIMMN, DTIMMX; Optional, required only if automatic time-step calculation is being used and calculation data are being read at this time.
  - DPTAS Maximum change in pressure allowed for setting the time step automatically  $(F/L^2)$ . Default set at 5 × 10<sup>4</sup> Pa.
  - DTTAS Maximum change in temperature allowed for setting the time step automatically (T). Default set at 5 °C.
  - DCTAS Maximum change in mass fraction (or scaled-mass fraction) allowed for setting the time step automatically (-). Default set at 0.25 (scaled).
  - DTIMMN Minimum time step required (t). This time step will be used for the first two steps after a change in boundary conditions, that is, at TIMCHG. Default set at 10<sup>4</sup> s.

DTIMMX - Maximum time step allowed (t). Default set at  $10^7$  s.

3.7.4 TIMCHG.

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TIMCHG - Time at which new transient data will be read or at which the simulation will be terminated (t).

Output information

3.8.1 PRIVEL, PRIDV, PRISLM, PRIKD, PRIPTC, PRIGFB, PRIWEL, PRIBCF.

PRIVEL - Printout interval (integer) for velocity arrays. These are interstitial velocities at the cell boundaries.

- PRIDV Printout interval (integer) for fluid density and fluidviscosity arrays.
- PRISLM Printout interval (integer) for solution-method information, number of iterations, maximum changes in dependent variables, and so forth.
- PRIKD Printout interval (integer) for conductance and dispersioncoefficient arrays.
- PRIPTC Printout interval (integer) for pressure, temperature, and mass-fraction arrays.
- PRIGFB Printout interval (integer) for flow-balance information for the region.
- PRIWEL Printout interval (integer) for well information.
- PRIBCF Printout interval (integer) for specified-value boundarycondition flow rates.
- For all of the above printout intervals:
  - 0 Means no printout of this information.
  - n Means that printout will occur every nth time step and at the end of the simulation.
  - -1 Means that printout will occur only at the time at which new transient data will be read and at the end of the simulation.
- 3.8.2 IPRPTC; Optional, required only if dependent-variable printouts are desired.
  - IPRPTC Index for printout of pressure, temperature and mass-fraction arrays. It is of the form  $n_1n_2n_3$  where  $n_1$  is for the pressure;  $n_2$  is for the temperature; and  $n_3$  is for the mass-fraction array. The  $n_i$  are set to 1 if printout is desired for the ith variable.  $n_1$  is set to 2 if both pressures and potentiometric heads are to be printed for isothermal cases.  $n_2$  is set to 2 if both temperatures and fluid enthalpies are to be printed.

3.8.3 CHKPTD (T/F), NTSCHK, SAVLDO (T/F).

CHKPTD - True if check-point dumps are to be made for possible restarts of the simulation.

NTSCHK - Number of time steps between successive check-point dumps. If set to -1, a dump will occur only at the times when new transient data are read and at the end of the simulation. SAVLDO - True if only the last check-point dump is to be saved.

The following four records are for the generation of contour maps on the line printer.

3.9.1 RDMPDT (T/F), PRTMPD (T/F).

RDMPDT - True if control data for map generation are to be read at this time.

PRTMPD - True if control data for map generation are to be written to the output file.

3.9.2 MAPPTC, PRIMAP; Optional, required only if contour-map-control data are to be read at this time.

MAPPTC - Index for a zoned contour map. It is in the form  $n_1n_2n_3$ , where  $n_1$  is for pressure;  $n_2$  is for temperature;  $n_3$  is for mass fraction. The  $n_i$  are set to 1 if a contour map is desired for the ith dependent variable.

PRIMAP - Printout interval (integer) for contour maps. Number of time steps between map generations.

0 - means no contour maps.

n - means contour maps at every nth time step.

-1 - means contour maps at the time when new transient data will be read and at the end of the simulation.

3.9.3 YPOSUP (T/F), ZPOSUP (T/F), LENAX, LENAY, LENAZ; Optional, required only if contour-map-control data are to be read at this time.

YPOSUP	-	True	if	the	y-axis	is	posit	tive	upwar	rd on	the	page	for	thi	S
		conto	our-	map	set.										
ZPOSUP	-	True	if	the	z-axis	is	posit	tive	upwar	d on	the	page	for	thi	s
		conto	our-	map	set.										
LENAX	-	Length	n of	the	x-axis	s on	the	page	for	this	cont	tour-r	nap	set	(in).
LENAY	-	Length	a of	the	y-axis	s 00	the	page	for	this	cont	tour-r	nap	set	(in).
LENAZ	-	Length	n of	the	z-axis	s on	the	page	for	this	cont	tour-	nap	set	(in).

- 3.9.4 IMAP1, IMAP2, JMAP1, JMAP2, KMAP1, KMAP2, AMIN, AMAX, NMPZON; Optional, required only if contour-map-control data are to be read at this time.
  - IMAP1, IMAP2 Range of node numbers along the x-axis for a contour map. Default set from 1 to NX.
  - JMAP1, JMAP2 Range of node numbers along the y-axis for a contour map. Set to 1,1 for cylindrical coordinates. Default set from 1 to NY.
  - KMAP1, KMAP2 Range of node numbers along the z-axis for a contour map. Default set from 1 to NZ.
  - AMIN, AMAX Range of the dependent variable for a contour map (appropriate units). If a pair of null entries, automatic scaling of the range will be performed of that dependent variable.

One pair of the indices iMAP1 and iMAP2 may be set equal to produce a contour map for just one plane.

NMPZON - Number of zones into which the contour map will be divided. Default set at 5. Limit of 32.

Up to three records of type 3.9.4 may be needed, depending on which combinations of pressure, temperature, and mass fraction are selected for mapping. The record order is: (1) Pressure-map data, (2) temperature-map data, and (3) solute-mass-fraction map data.

This ends the transient data set that is read at a given time. At simulation time equal to TIMCHG, another transient-data set will be read, until the simulation is finished. At that time, THRU is read as true in the following record.

3.99.1 THRU (T/F).

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THRU - Set to true at this point to signify the end of the simulation.

3.99.2 PLOTWP(T/F), PLOTWT(T/F) PLOTWC(T/F).

PLOTWP - True if observed and(or) calculated well pressures are to be plotted versus time.

PLOTWT - True if observed and(or) calculated well temperatures are to be plotted versus time.

PLOTWC - True if observed and(or) calculated well mass fractions are to be plotted versus time.

Temporal-plot information

The following data records of type 4.N are required only if character-string plots of variables versus time are desired at selected wells.

- 4.1 IWEL, RDPLTP (T/F); Optional, required only if temporal plots are to be made, and new plot-control parameters are to be set for subsequent plots (RDPLTP is true).
  - IWEL Well number. This number must agree with the number associated with the calculated data.

RDPLTP - True if new plotting-control parameters are to be read at this time for subsequent plots.

4.2 IDLAB; Optional, required only if temporal plots are to be made, and new plot-control parameters are to be read for subsequent plots (RDPLTP is true).

- IDLAB Identification label for this well's plots. A character string of up to 80 characters. Space over at least one character position from the left.
- 4.3 NTHPTO, NTHPTC, PWMIN, PWMAX, PSMIN, PSMAX, TWMIN, TWMAX, TSMIN, TSMAX, CMIN, CMAX; Optional, required only if temporal plots are to be made, and new plot-control parameters are to be read for subsequent plots (RDPLTP is true).
  - NTHPTO Index for plotting the first, then every nth observed data point versus time. Default set to one. A blank may be entered if no observed data are to be plotted.
  - NTHPTC Index for plotting the first then every nth calculated value versus time. Default set to one.
  - PWMIN, PWMAX Minimum and maximum values of pressure at the well datum that set the axis range for the temporal plots  $(F/L^2)$ .
  - PSMIN, PSMAX Minimum and maximum values of pressure at the land surface that set the axis range for the temporal plots  $(F/L^2)$ .
  - TWMIN, TWMAX Minimum and maximum values of temperature at the well datum that set the axis range for the temporal plots (T).
  - TSMIN, TSMAX Minimum and maximum values of temperature at the land surface that set the axis range for the temporal plots (T).
  - CMIN, CMAX Minimum and maximum values of solute-mass fraction (scaledmass fraction) at the well datum and the land surface that set the range for the temporal plots (-).

The pressure, temperature and solute-mass fraction ranges for the plots can be specified by the user or established automatically. The latter option is invoked by entering zeros for the maximum and minimum values. 4.4 TO, POW, POS, TOW, TOS, COW; Optional, required only if there are observed data.

TO - Time of observation (t).

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- POW Pressure observed at the well-datum level  $(F/L^2)$ .
- POS Pressure observed at the land surface in the well  $(F/L^2)$ .
- TOW Temperature observed at the well-datum level (T).
- TOS Temperature observed at the land surface in the well (T).

COW - Mass fraction (or scaled-mass fraction) observed in the well at the well datum or the land surface (-).

It is assumed that the observed data are in the same units that will be used for output of the calculated data. As many records of type 4.4 are used as necessary to enter all the observed well data. There may be wells for which only calculated data are available; for these wells, no records of type 4.4 will be read.

4.5 End this data set with -1. / .

Indicates the end of the observed data set for this well.

As many records of type 4.1-4.4 are used as necessary for all of the wells for which observed or calculated data are being plotted.

4.6 End this data set with 0 / .

Indicates the end of the temporal-plot information and the observed data for all the wells.

This ends the input-data-file description. For quick reference, a list of the definitions for the various program-control options is provided in table 5.3.

Variable		Option definitions
SLMETH	1 -	Selects the direct, D4, equation solver.
		Selects the iterative, two-line, successive-
	-	overrelaxation equation solver.
the follow:	ing, n <sub>1</sub> den	otes outward normal direction to the boundary face
		is the y-direction; and 3 is the z-direction.
IBC		Cell is excluded from the simulation region.
		Specified-pressure boundary-condition node.
	010 -	Specified-temperature boundary-condition node.
		Specified-solute-concentration boundary-condition node.
		Specified-fluid-flux boundary-condition cell.
		Specified-diffusive-heat-flux boundary-condition
	-	cell.
	n <sub>1</sub> 00002 -	Specified-diffusive-solute-flux boundary-condition cell.
		Leakage boundary-condition cell.
		Aquifer-influence-function boundary-condition cell
		Heat-conduction boundary-condition cell.
	1100040	
IAIF	1 -	Pot aquifer for outer region.
		Transient-aquifer-influence function with
		calculation using the Carter-Tracy approximation.
WQMETH	10 -	Specified well-flow rate with allocation by
		mobility and pressure difference.
		Specified well-flow rate with allocation by mobility.
		Specified pressure at well datum with allocation
		by mobility and pressure difference.
		Specified well-flow rate with a limiting pressure
		at well datum. Flow-rate allocation by mobility
		and pressure difference.
	40 -	Specified surface pressure with allocation by
		mobility and pressure difference. Well-riser
		calculations will be performed.
		Specified surface-flow rate with limiting surface
		pressure. Allocation by mobility and pressure
		difference. Well-riser calculations will be
		performed.
		Observation well or abandoned well.
FDSMTH	0.5 -	Centered-in-space differencing for advective terms
		Upstream differencing for advective terms.
FDTMTH	0.5 -	Centered-in-time differencing.

Table 5.3.--Option lists for program-control variables

Variable	Option definitions						
IPRPTC	lxx - Printout of pressure field.						
	2xx - Printout of pressure and potentiometric-head fields						
×	xlx - Printout of temperature field.						
	x2x - Printout of temperature- and fluid-enthalpy fields.						
	xx1 - Printout of solute-concentration field.						
ORENPR	12 - Printouts of arrays by areal (x-y) layers.						
	13 - Printouts of arrays by vertical x-z or (r-z) slices						
A neg pag	ative value means the y or z-axis is to be positive down the						
-	ative value means the y or z-axis is to be positive down the e.						
pag	ative value means the y or z-axis is to be positive down the						
pag	ative value means the y or z-axis is to be positive down the e. 0 - No printout. n - Printout every nth time step and at the end of the						
pag	<pre>ative value means the y or z-axis is to be positive down the e. 0 - No printout. n - Printout every nth time step and at the end of the simulation.</pre>						
pag	<ul> <li>ative value means the y or z-axis is to be positive down the e.</li> <li>0 - No printout.</li> <li>n - Printout every nth time step and at the end of the simulation.</li> <li>-1 - Printout only at the time of new transient data</li> </ul>						
pag PRIxxx	<ul> <li>ative value means the y or z-axis is to be positive down the e.</li> <li>0 - No printout.</li> <li>n - Printout every nth time step and at the end of the simulation.</li> <li>-1 - Printout only at the time of new transient data being read and at the end of the simulation.</li> </ul>						

Table 5.3.--Option lists for program-control variables--Continued

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# 6. OUTPUT DESCRIPTION

Various types of output result from running the HST3D program. Most of the output is to disc files to be displayed on a video screen or routed to a line printer. These files are written in ASCII format. The two exceptions are the optional check-point/restart dumps written in binary format to a disc file, and the calculated dependent-variable data for the wells that periodically are written, also in binary format, to a disc file for the temporal plots that can be made at the end of the simulation.

Output is generated at several stages during the simulation. Some information, such as the heading, title, array-partitioning data, and problem-geometrical information, is printed always. The heading contains the program version number which will change when major modifications or corrections are made. The units employed for the ouput are the same as those used for the input data, either metric or inch-pound as specified in record 1.4. Table 5.2 and the input-record descriptions (section 5.2.2) give the inch-pound and the metric units employed. For easier reading, variables are identified in the output by descriptors rather than program-variable names. Much of the output is optional, and the numbers of the records containing the control variables in the data-input-form list of table 5.1 are indicated. The writing of a file that echos each record of input data, as it is read, is optional (record 1.10). The static data that may be printed include porous-media properties, fluid properties, initial-condition distributions, boundarycondition information, solution-method information, well information (record 2.23.1), and density and viscosity distributions (record 2.23.2). The selection of which of the dependent variables (pressure, temperature or mass fraction) will have initial conditions printed is made in record 2.23.2.

Print intervals can be selected individually for information that is printed at the end of a time step. The information printed may include the velocity distribution, the density and viscosity distributions, the solution method information, the conductance and dispersion-coefficient distributions, the dependent-variable distributions, the regional fluid-flow, heat-flow and solute-flow rates, the regional cumulative-flow results, and the specified-

value boundary-condition flow rates (record 3.8.1). The selection of of the dependent variables which will be printed is determined in record 3.8.2.

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Contour maps of pressure, temperature, and mass fraction can be produced on the line printer; they are zoned into intervals and may cover subregions of the simulation as specified by the user (record 3.9.4). The contour-mapping routine produces character-string plots. Alternating zones of symbols and blanks are used to make perception easier. The user can make a programming change (set variable, ZEBRA, to false) to cause the symbol-filled zones to be adjacent to each other. Contour intervals are automatically calculated to be a multiple of 2, 5, or 10. The lower and upper limits can be chosen by the user or determined from the range of the data to be contoured. In the former case, values below the specified-lower limit are contoured with a zone of minus signs and values above the specified-upper limit are contoured with a zone of plus signs. The contour zones contain their lower-boundary values: the upper-boundary values belong to the next zone above with the exception of ~ the highest zone of the map which does contain its upper-boundary value. The maps are either areal or vertical slices along nodal planes of the threedimensional region (record 2.23.3), with the orientation sepcified in record 3.9.3. The size of the maps on the paper is chosen by the user (record 3.9.3). An echo printout of the mapping specifications can be requested (record 3.9.1). Bilinear interpolation is used to locate the contour-interval boundaries; cells excluded from the simulation region are indicated by X's. If multiple pages are used for the contour maps, no printing is done across the paper folds. Thus, separation and alinement of the various pages is necessary to eliminate gaps.

Temporal plots of selected variables are also in character-string format. The plots that may be produced at the end of the simulation include well-datum pressure, well-surface pressure, well-datum temperature, well-surface temperature, and well solute-mass fraction (or scaled-mass fraction) (record 3.99.2). For observation wells, the well-datum value is taken to be the value in the aquifer cell at the well-datum level. Observed (record 4.4) and calculated data of the same type are plotted together for comparison purposes. The time axis runs down, and the dependent-variable axis runs across the page.

A limit of 500 lines is set. If the time series to be plotted of any calculated variable contains more than three times the total number of node points in the region, or the series of any observed variable contains more than two times the number of node points in the region, array-storage problems will occur and program execution will be terminated. These problems may be avoided by plotting the first point followed by only every nth point thereafter (record 4.3). The user may specify the ranges of the variables to be plotted. However automatic scaling of the plot is available using the minimum and maximum values of the variables. Axis subdivisions that are a multiple of 2, 5 or 10 are produced. The present version of the HST3D code contains no provision for producing line plots on pen-plotting devices or video screens.

## 7. COMPUTER-SYSTEM CONSIDERATIONS

The heat- and solute-transport simulation program was developed initially on a Control Data Cyber 170/720 computer<sup>1</sup>, and finally on a Prime 9950 computer. The Cyber computer has a very fast arithmetic central-processing unit relative to the Prime, while the Prime has virtual storage that the Cyber does not have. Therefore, the programming philosophies needed to create the optimum code for execution of large, long-running simulations are in direct opposition for these two machines. Specifically, the Cyber, with its fast arithmetic, but limitations on storage, is most efficiently used with a code that minimizes storage requirements. This is accomplished to a certain extent by recalculating some quantities each time they are needed, rather than storing them. On the other hand, the virtual storage of the Prime means that the running time for large, long simulations may become inconveniently long. This implies that the most efficient code for the Prime will use more storage than the Cyber and never compute a quantity more than once.

The present version of the heat- and solute-transport code is not optimal for either type of machine, but it tends to be oriented toward the Prime. Further optimization of the program will require timing tests. The storage requirement on the Prime computer for the executable-code module is about 1.1 megabytes, exclusive of the variably partitioned arrays, when compiled with the interactive-debug option and no optimization.

The language used for this program is FORTRAN-77, although some FORTRAN-IV coding still exists. An attempt has been made to use only the ANSI standard FORTRAN-77 for maximum portability (American National Standards Institute, 1978).

Double-precision arithmetic has been used for all real variables. Separate variably partitioned arrays were defined for real and integer vari-

<sup>1</sup>Use of brand names in this report is for identification purposes only and does not constitute endorsement by the U.S. Geological Survey.

ables, so that the real variables could be made double precision. FORTRAN-77 intrinsic function names were used in their generic form, so that no changes had to be made for use with double-precision variables.

Although the standards for FORTRAN-77 have been followed as closely as possible, there are always problems of portability to different computer systems. Development experiences revealed some differences between the Prime and Cyber computers that affect program portability.

There are no alternate entry points into any of the subroutines, to avoid computer incompatibilities. The Prime uses dynamic storage of variables and of compiled code during program execution. However, all variables in common blocks are automatically made static. Static data items retain their values between subprogram references, while dynamic data items in a subprogram lose their values upon return from that subprogram. By including all common blocks in the main program, potential problems with computer systems that do not make common block variables automatically static should be avoided. Variables are explicitly initialized where necessary, so no reliance is made on systemdefault initialization.

The Cyber computer FORTRAN compiler allows only 63 arguments in a subprogram-argument list. The Prime compiler allows 256. There are a few subroutines in the heat- and solute-transport program that have more than 63 arguments. To reduce the number to within the limit for the Cyber computer, some subarrays would need to be eliminated from the argument list, necessitating some recoding. The eliminated subarrays would be passed by passing the entire variably partitioned arrays along with appropriate pointer indices.

The Cyber requires the BLOCKDATA subprogram to be contained in the same file as an executable subprogram; whereas, the Prime allows it to be compiled from its own separate file. The version of the code discussed in this documentation has the main program, and each subprogram, including BLOCKDATA, contained in a separate file.

#### 8. COMPUTER-CODE VERIFICATION

Verification of a computer program is the process of ensuring that the code performs the intended calculations correctly. This is in contrast to computer-model validation, which is the demonstration that a particular model with a particular set of parameters adequately describes a given physical situation. Program verification is accomplished by running various test problems for which an analytical solution is known, or for which numerical results from another verified program are available.

#### 8.1. SUMMARY OF VERIFICATION TEST PROBLEMS

Several sets of test problems have been used for verification and are summarized hereinafter. Verification is a continuing process, as many combinations of program options could be tested.

## 8.1.1. One-Dimensional Flow

Test problem set 1 tested the ability of HST3D to simulate compressible flow and was based on the analog between confined ground-water flow and heat conduction. A thermal-conduction problem was taken from Carnahan and others (1969, p. 443). The physical situation was that of one-dimensional confined flow of a compressible fluid when a unit-step increase of pressure was applied at both ends of the region. The dimensions of the simulation region were 1.0 meter in each direction. Eleven equally spaced nodes were used to discretize the region in the x-direction, with two nodes each in the y- and z-directions. The initial condition was hydrostatic equilibrium. The porosity was set to 1.0 so no porous medium was present. The parameters were set to the following values:

Porosity, 1.0 Permeabilities

x-direction,  $1.\times10^{-8}$  m<sup>2</sup>; y- and z-directions  $1.\times10^{-20}$  m<sup>2</sup>

Fluid compressibility, 1×10<sup>-5</sup> Pa<sup>-1</sup>

Fluid density (at reference conditions), 1,000 kg/m<sup>3</sup>

Fluid viscosity (at reference conditions), 0.001 kg/m-s. A fixed time step of  $1.25 \times 10^{-2}$  was used and the results compared at  $2.5 \times 10^{-2}$ s. Backwards-in-time differencing was used with the direct method for matrix

solution.

Variations on the basic problem included using centered-in-time differencing, reduction of the time step length by a factor of 10, using the iterative solver for the matrix equation, and entering data in inch-pound units.

The results agreed to five significant digits with the numerical solution of Carnahan and others (1969, p. 446-447). Representative values for pressure increase at  $2.5 \times 10^{-2}$ s are shown in table 8.1. The results were symmetric about the mid-point of the region in the x-direction as expected.

Distance along column	Change in pressure calculated	Change in pressure calculated
(meters)	by HST3D (Pa)	by Carnahan (Pa)
0	1.00000	1.00000
0.2	0.32471	0.32471
0.4	0.10104	0.10104
0.5	0.07965	0.07965

Table 8.1.--Representative values for the pressure increase from the HST3D simulator and the results of Carnahan and others (1969) [Values at time of  $2.5 \times 10^{-2}$  seconds]

### 8.1.2. Flow to a Well

Flow to a single well in a cylindrical-coordinate system was the basis for test problem set 2, providing another test of the flow-simulation part of HST3D. Both confined and unconfined conditions were simulated. The confined-flow problem was taken from Lohman (1972, p. 19) and the unconfinedflow problem was example 1 from Boulton (1954). A specified flow rate from the well was used for both problems.

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A cylindrical region of 2,000 ft external radius and 100 ft thickness was used for the confined-flow problem. The theoretical results of Lohman were based on the this solution for an infinite region, so comparisons were restricted to time values sufficiently small so that the outer boundary did not affect the flow field. Twenty-one nodes were used in the radial direction logarithmically spaced by the automatic discretization algorithm, except for those at 200 and 400 ft. Two nodes defined the vertical discretization. The well flow rate was allocated by mobility and the upper, lower and outer boundaries were impermeable. The initial condition was that of hydrostatic equilibrium. A time step of 30 s was used for a duration of 600 s. The parameters used for the confined problem were the following:

Porosity, 0.20 Fluid compressibility, 3.33×10<sup>-6</sup> psi<sup>-1</sup> Porous-medium compressibility, 3.94×10<sup>-6</sup> psi<sup>-1</sup> Permeability, 5.31×10<sup>-10</sup> ft<sup>2</sup> (hydraulic conductivity 137 ft/d) Well radius, 0.1 ft Well flow rate, 96,000 ft<sup>3</sup>/d Fluid density at reference conditions, 62.4 lb/ft<sup>3</sup> Fluid viscosity at reference conditions, 1 cP

The calculated fluid drawdown was compared with the results of Lohman (1972, p. 19) for several locations at six values of time. The drawdown values agreed to within 0.01 to 0.1 ft (table 8.2). The differences were due mostly to spatial-discretization error, since they were reduced by 30 to 50 percent by doubling the number of nodes in the radial direction.

A cylindrical region of 2,000 ft radius and 800 ft thickness was used for the unconfined-flow problem. The theoretical solution of Boulton (1954) was used for comparison. This solution was based on a linearized free-surface boundary condition valid for small values of drawdown and included a correction for the fact that a line sink of constant intensity represented the well flow.

Sixteen nodes were distributed in the radial direction, logarithmically spaced, except at 160 ft, and nine nodes were distributed in the vertical direction. Advantage was taken of the fact that the free surface can rise above the upper plane of nodes. The well flow rate was allocated by mobility. The lower and outer boundaries were impermeable while the upper boundary was a free surface. The initial condition was that of hydrostatic equilibrium. The automatic-time-step algorithm was used to simulate from 10 s to  $3 \times 10^5$  s.

	Drawdown (feet)						
		(feet)	et)				
Time (second)		200	400				
	HST3D	Lohman	<b>HST3D</b>	Lohman			
60	0.56	0.66	0.15	0.16			
120	0.91	0.99	0.33	0.38			
240	1.29	1.36	0.61	0.67			
300	1.42	1.49	0.72	0.77			
480	1.67	1.75	0.94	0.99			
600	1.79	1.86	1.05	1.12			

Table 8.2.--Comparison of drawdown values calculated by HST3D with those of Lohman (1972, p. 19)

The parameters for the unconfined-flow problem were the following:

Porosity, 0.15 Fluid compressibility, 1×10<sup>-15</sup> psi<sup>-1</sup> Porous-medium compressibility, 1×10<sup>-15</sup> psi<sup>-1</sup> Permeability, 6.7×10<sup>-12</sup> ft<sup>2</sup> (hydraulic conductivity 2×10<sup>-5</sup> ft/s) Well radius, 0.1 ft Well flow rate, 1 ft<sup>3</sup>/s Fluid density at reference conditions, 62.4 lb/ft<sup>3</sup> Fluid viscosity at reference conditions, 1 cP

The calculated drawdown of fluid was compared with the result of Boulton's (1954) example 1. His only reported value was at a time of 3.47 days and a radius of 160 ft. The drawdown was 2.13 ft compared to the HST3D result of 2.20 ft. Use of five nodes in the vertical direction reduced the drawdown calculated by HST3D to 1.96 ft. This indicated the effect of spatial-discretization error, particularly when vertical flow is important.

The agreement is very good considering that the HST3D simulator does not take the kinematic boundary condition at the free surface into account. Therefore greater discrepancies should appear as the well bore is approached.

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### 8.1.3. One-Dimensional Flow

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Three cases of ground-water flow for which analytic solutions are available (Bear, 1972, p. 301, 367, 380) formed test problem set 3. They were one-dimensional, confined flow; one-dimensional, unconfined flow; and one-dimensional, unconfined flow with influx from precipitation. The simulation region was  $400 \times 400 \times 100$  meters in the x-, y- and z-directions respectively. The flow field was horizontal in the y-direction for all cases. Specified-pressure boundary conditions were used on the inlet and outlet boundaries with impermeable lateral and bottom boundaries.

The parameters employed were the following:

Porosity, 0.15

Fluid compressibility, 5×10<sup>-15</sup> Pa<sup>-1</sup>

Porous-medium compressibility, 8.8×10<sup>-14</sup> Pa<sup>-1</sup>

Permeability, 1.18×10<sup>-11</sup> m<sup>2</sup>

Fluid density at reference conditions, 1,000 kg/m<sup>3</sup> Fluid viscosity at reference conditions, 0.001 kg/m-s

Spatial discretization was accomplished using five equally spaced nodes in the x- and y-directions and two nodes in the z-direction. All cases were run to steady-state with a time step of 86,400 s.

Case 1: Confined flow

The upper boundary surface was made impermeable and the other boundary conditions for this case were specified potentiometric heads of 200 m along the y = 0 boundary and 100 m along the y = 400 m boundary. The analytical solution was a linear potentiometric-head variation between the two boundaries (Bear, 1972, p. 301). The results at steady-state agreed with the analytical solution to five significant digits. Global-balance results were verified by hand calculation.

Case 2: Unconfined flow

The upper boundary was defined as a free surface and the region dimension in the z-direction was extended to 200 m. The permeabilities in the x-, y-, and z-directions were modified to  $1.18 \times 10^{-9} \text{ m}^2$ ,  $1.18 \times 10^{-10} \text{ m}^2$  and  $1.13 \times 10^{-5} \text{ m}^2$ respectively. The high permeability in the z-direction was to make the Dupuit approximation of hydrostatic equilibrium in the vertical direction valid. The specified potentiometric heads were 200 m along the y = 0 m boundary and 150 m along the y = 400 m boundary.

The simulation was run to 172,800 s, which was essentially steady-state. The results for potentiometric head were compared with the analytical solution (Bear, 1972, p. 367) based on the Dupuit approximation. Agreement to five significant digits was obtained at the three interior-node locations along the y-axis. The global-balance results were verified by hand calculation.

Better agreement with the analytical solution was obtained for this test problem than for the unconfined flow to a well in test problem set 2. This improved agreement was attributed to the fact that the analytical solution for this problem was based on the Dupuit assumption of negligible vertical flow and a high vertical-permeability value was used in the HST3D simulation to achieve hydrostatic equilibrium. The unconfined case of test problem set 2 had significant vertical flow. Because the HST3D simulation does not attempt to satisfy the non-linear, kinematic, free-surface boundary condition, described in sections 2.5.6 and 3.4.6, the poor results obtained in cases of significant vertical flow at the free surface were not surprising.

Case 3: Unconfined flow with precipitation recharge

Case 2 was modified by the addition of areal recharge at a uniform rate and distribution. The region dimension in the z-direction was extended to 275 m. The areal-recharge flux was set to  $-1,157 \times 10^{-3} \text{ m}^3/\text{m}^2\text{s}$  with the negative sign denoting flux in the negative z-direction.

The simulation was run to 172,800 s, which was essentially steady-state. The results for potentiometric head were compared with the analytical solution from Bear (1972, p. 380). This solution also was based on the Dupuit approximation. Agreement to five significant digits was obtained at the three interior-node locations along the y-axis. The global-balance results were verified by hand calculation.

### 8.1.4. One-Dimensional Solute Transport

Flow with solute transport in a one-dimensional column was the basis of test problem set 4. A steady-state flow field was established by specifying an initial-pressure gradient along the column. The boundary condition at the column inlet was a specified scaled-solute concentration of a unit-step at time zero. The column length was 160 m discretized by 21 equally distributed nodes in the x-direction. The y- and z-directions were 1 m with two nodes in each direction. The following parameter values were chosen:

Porosity, 0.5

Fluid compressibility, 1×10<sup>-10</sup> Pa<sup>-1</sup>

Porous-medium compressibility, 1×10<sup>-10</sup> Pa<sup>-1</sup>

Permeability, 1×10<sup>-10</sup> m<sup>2</sup>

Fluid density at reference conditions, 1,000 kg/m<sup>3</sup> (independent of solute concentration)

Fluid viscosity at reference conditions, 0.001 kg/m-s Interstitial velocity along the column,  $2.7778 \times 10^{-4}$  m/s Longitudinal dispersivity, 10 m Molecular diffusivity,  $1 \times 10^{-10}$  m<sup>2</sup>/s

The initial solute concentration in the column was zero. A time-step length of 720 s was used for a total simulation time of 7,200 s. The HST3D options tested included the spatial and temporal differencing methods and the two different equation solvers. A second case with a column that was 4-meters wide was also tested.

The results were compared to a one-dimensional finite-difference transport program (Grove and Stollenwerk, 1984), and to an analytical solution (Ogata and Banks, 1961). The calculated solute-mass fraction values agreed with the finite-difference program results to four significant digits. Differences between the HST3D results and the analytical solution were as much as 0.035 units of scaled-mass fraction. The results at the end of the simulation period appear in table 8.3.

Table 8.3.--Scaled solute-concentration values calculated by HST3D compared to the one-dimensional finite-difference solution of Grove and Stollenwerk (1984) and the analytical solution of Ogata and Banks (1961)

<sup>[</sup>Values at time of 7,200 seconds; CSCT, centered-in-space and centered-intime differencing; BSBT, backward-in-space and backward-in-time differencing]

,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			يعكيد المتبعد المنتي ككانك	solute conc ance along (meters		)
		0	8	16	24	32
HST3D	CSCT BSBT	1.0000	0.31665 0.37500	0.05939 0.09414	0.007843	0.000801
One-dimensional finite-differ- ence solution	CSCT	1.0000	0.31666	0.05939	0.007843	0.000801
Analytical solution		1.0000	0.29808	0.02439	0.000469	0.000002

The discrepancies are attributed to spatial- and temporal-discretization errors, because reducing the spatial and temporal steps by factors of eight and five respectively, reduced the maximum difference to 0.003 units of scaled-mass fraction. The differences between the two differencing schemes were the result of numerical-dispersion errors. The simulation was not run long enough for the difference between the analytic and numerical boundary condition at the far end of the column to affect the solution. Flow and solute global-balance residuals were at least eleven orders of magnitude smaller than the net amounts entering the region.

## 8.1.5. One-Dimensional Heat Conduction

Test problem set 5 involved heat transport without fluid flow. A heatconduction problem was taken from Carnahan and others (1969, p. 443). The physical situation was that of one-dimensional heat conduction when a unitstep increase of temperature was applied at both ends of a column. The dimensions of the region were 1 m in each direction. Eleven equally spaced nodes were used to discretize the region in the x-direction, with two nodes each in the y- and z-directions. The initial condition was a uniform temperature of 1 °C. The parameters were set to the following values:

Porosity, 1.0 (no porous medium present) Fluid compressibility, 5×10<sup>-6</sup> Pa<sup>-1</sup> Permeability, 1×10<sup>-8</sup> m<sup>2</sup> Fluid density, 1,000 kg/m<sup>3</sup> Fluid viscosity, 0.001 kg/m-s (independent of temperature) Fluid thermal expansion factor, 0. °C<sup>-1</sup> Fluid heat capacity, 1.0 J/kg °C Fluid thermal conductivity, 1.0 W/m-°C

A time step of 0.0125 s was used for a total simulation time of 0.0250 s. The options tested included backwards-in-time differencing, direct and iterative solvers of the matrix equations, and inch-pound and metric units for data entry and output. The results agreed to five significant digits with those of Carnahan and others (1969, p. 446, 447) and matched the numerical values for scaled-pressure rise given in table 8.1. Heat-balance residuals were 12 orders of magnitude less than the amount of heat that entered the region.

## 8.1.6. One-Dimensional Heat Transport

Test problem set 6 was heat transport with fluid flow and was the analog to problem s.t 4. A steady-state flow field was established by specifying an initial-pressure gradient along the column. The boundary condition at the column inlet was a specified scaled-temperature of a unit-step at time zero. The dimensional-temperature step was 10 °C. The column length was 160 m discretized by 21 equally distributed nodes in the x-direction. The y- and z-directions were 1 m with two nodes in each direction. The following parameter values were used:

Porosity, 0.5
Fluid compressibility, 1x10<sup>-10</sup> Pa<sup>-1</sup>
Porous-medium compressibility, 1x10<sup>-10</sup> Pa<sup>-1</sup>
Permeability, 1x10<sup>-10</sup> m<sup>2</sup>
Fluid density at reference conditions, 1,000 kg/m<sup>3</sup> (independent of temperature)
Fluid viscosity at reference conditions, 0.001 kg/m-s (independent of temperature)
Interstitial velocity along the column, 2.7778x10<sup>-2</sup> m/s
Longitudinal dispersivity, 10 m
Porous-medium product of density and heat capacity, 800 J/m<sup>3</sup>-°C
Porous-medium thermal conductivity, 1.8 W/m-°C
Fluid heat capacity, 4,200 J/kg-°C
Fluid thermal conductivity, 0.6 W/m-°C
Fluid thermal expansion factor, 0. °C<sup>-1</sup>

The initial temperature in the column was 10 °C. A time step length of 1076.5 s was used for a total simulation time of 10,765 s. Centered-in-space and centered-in-time differencing were used for discretization.

The results were compared to a one-dimensional finite-difference transport program (Grove and Stollenwerk, 1984), and to an analytical solution (Ogata and Banks, 1961). The scaled temperature values matched the analogous solute-transport problem of set 4 as expected. Numerical differences in the fourth significant digit were attributed to the fact that the thermaldispersion coefficient was slightly larger than the solute-dispersion coefficient. The results at the end of the simulation period appear in table 8.4.

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Table 8.4.--Scaled temperature values calculated by HST3D compared to the one-dimensional finite-difference solution of Grove and Stollenwerk (1984) and the analytical solution of Ogata and Banks (1961)

	Scaled temperature (-) Distance along the column (meters)									
	0	8	16	24	32	56				
HST3D	1.0000	0.31670	0.05941	0.007846	0.000802	0.000005				
One-dimen- sional finite- difference		•								
solution	1.0000	0.31671	0.05941	0.007847	0.000802	0.00000				
Analytical solution	1.0000	0.29815	0.02441	0.000470	0.000002	0.00000				

[Values at time of 10,765 seconds]

The time at which the temperature profile essentially matched the soluteconcentration profile is a factor of about 1.5 later. This is because the thermal-storage coefficient, which includes the porous-matrix solid phase as well as the fluid phase, is greater than the solute-storage coefficient involving only the fluid phase. The effect of spatial- and temporaldiscretization errors can be seen by comparing the results to the analytical solution. The simulation was not run long enough for the difference between the analytic and the numerical boundary condition at the far end of the column to affect the solution. Flow and heat global-balance residuals were at least eight orders of magnitude smaller than the net amounts entering the region.

### 8.1.7 Thermal Injection in a Cylindrical Coordinate System

Simulation of the injection of hot water at 60 °C into an aquifer initially at 20 °C for 90 days, followed by production for an equivalent time period, formed test problem set 7. A cylindrical-coordinate system was used with a fully penetrating well. Both density and viscosity were taken to be functions of temperature. Results from this test problem were compared to the results obtained by Voss (1984) p. 207-212 using his SUTRA finite-element transport-simulation program. Options tested included the approximate, augment.d-diagonal treatment of the cross-derivative dispersion fluxes, the explicit evaluation of the cross-derivative dispersion fluxes, central and upstream differencing for the advective terms, and equal and unequal longitudinal and transverse dispersivities.

A list of the parameters employed will not be presented here, because this test problem also is given as an example for the user. The complete set of parameters and other data that define the problem is presented in section 8.2.2.

The region had an interior radius of 1 m, exterior radius of 226 m, and a thickness of 30 m. At the upper and lower surfaces, the boundary conditions were no fluid flow and no heat flow. At the exterior radius, a hydrostatic pressure was specified with any fluid entering the region having a temperature of 20 °C. The initial condition was hydrostatic equilibrium of the fluid at 20 °C.

The results for the temperature field at 30, 90, 120, 150, 180 days were compared to those of Voss (1984, p. 207-212) for his particular case of options. The profiles were in general agreement with some deviation for the withdrawal phase. The warmer water was extracted more rapidly in the HST3D simulation. This can be attributed to the fact that Voss used a line sink of constant intensity per unit length to represent the well whereas the HST3D simulator allocated the flow from each layer in proportion to the local-fluid mobility. The increased mobility of the warmer fluid caused increased flow from the upper parts of the well bore and decreased flow from the lower. The

total-flow rate was held constant. The finite-difference methods used by the HST3D simulator showed some spatial oscillation of about 3 °C. Use of upstream differencing removed this problem at the cost of increased dispersion which reduced the sharpness of the temperature front considerably.

## 8.2. TWO EXAMPLE PROBLEMS

1.1.1

Two example problems are presented for the purpose of giving the new HST3D program user some experience in learning to run a successful simulation. One involves solute transport and the other involves heat transport. These examples also will aid in adapting the HST3D program to run on computer systems other than the PRIME. The problem descriptions, data files and selected parts of output are included for comparison. Several ways are available to input some of the data so an exact match with the data files presented is not necessary to execute these examples correctly.

8.2.1. Solute Transport with Variable Density and Variable Viscosity

The first example problem is based upon displacement of a fluid of one density and viscosity by another fluid of different density and viscosity. The density and viscosity differences are caused by different amounts of dissolved solute. The system is isothermal. The region is a square slice of porous medium, that is oriented vertically, so that gravitational effects will occur in the flow field. The dimensions are 2 m in the x-direction by 0.2 m in the y-direction by 2 m in the z-direction. The z-direction is oriented vertically upward. Except at the inlet and outlet corners, the boundaries are confining. The porous medium is homogeneous and isotropic.

The parameters to be used are as follows:

Permeability, 1×10<sup>-8</sup> m<sup>2</sup> Porosity, 0.10 Density of fluid initially present, 800 kg/m<sup>3</sup> Viscosity of fluid initially present,  $1.3 \times 10^{-3}$  kg/m-s Scaled-solute-mass fraction of fluid initially present, 0. Temperature of region (isothermal), 20 °C Density of injected fluid, 1,000 kg/m<sup>3</sup> Viscosity of injected fluid,  $7 \times 10^{-4}$  kg/m-s Scaled-solute-mass fraction of injected fluid, 1.0 Longitudinal dispersivity, 0.1 m Transverse dispersivity, 0.1 m Fluid compressibility, 0. Pa<sup>-1</sup>

The molecular diffusivity of the solute is neglected. Absolute mass fractions of solute are needed for solute-mass balance calculations. They may be chosen as 0.0 for the fluid initially in the region and 0.005 for the injected fluid. The initial condition is one of hydrostatic equilibrium with a pressure of 0.0 Pa at an elevation of 2 m. The injection location is at the lower left-hand corner of the region. The injection boundary is maintained at a scale-solute concentration of 1.0 with an injection pressure of 25,000 Pa. The outlet is at the upper right-hand corner of the region and is open to the atmosphere. The region is illustrated in figure 8.1.

Construct a numerical model of this system with nodal dimensions  $11 \times 2 \times 11$  and observe the migration of the fluid containing solute from the lower left-hand corner to the upper right-hand corner for a total simulation time of 10 s. Use a time-step length of 0.2 s. Use the approximate method for calculating the cross-derivative dispersive terms. Print out results at 10 s with a contour plot of solute concentration. Use a contour interval of 0.2.

A listing of the data file that will run example problem 1 is given in table 8.5. The input-data form (table 5.1) was used to construct this file, but comments pertaining to unnecessary data items have been eliminated for brevity.

The output file for this problem is contained in table 8.6. The header shows the release number for the version of the program. The problem title and information relating to dimensioning requirements is next. Then follows the static data. Read-echo printouts were selected for data input by i,j,k range. The conductance factors are constant for this simulation. The numbering sequence for boundary-condition cells is primarily for debugging purposes. In this problem, the approximate method for handling the cross-derivative dispersion terms using amplified-diagonal values was chosen.

The next section of the printout contains the transient data including both input parameters and output variables at selected time steps. For this example, the input data for boundary conditions, calculation information, and mapping data are printed. The printout interval was set to print at the end of the simulation only at time step 50. No cross-derivative dispersive conductances appear, because the approximate method was selected for handling cross-dispersive fluxes. The output at the end of the time step, and end of the simulation in this case, includes some calculation information, pressure and solute-mass fraction, density and viscosity, the global-balance summary, boundary-condition flow rates, a contour map of solute-mass fraction, and the velocity field. The contour map was made for only one plane of cells because the other is identical by symmetry. The velocity field is that which would be used for the next time step, if one were to be calculated. In the globalbalance summary, we can see that the flow-balance residual is about 5 orders of magnitude less than the other amounts. The solute-balance residual is about 5 orders of magnitude less than the amount of inflow and amount of change. Similar results exist for the cumulative amounts. The map of the solutemass fraction field shows the asymmetry that is caused by the denser injected fluid tending to stay in the bottom part of the region. The same case was run under conditions of constant density and viscosity and the results were symmetric about the diagonal as expected.

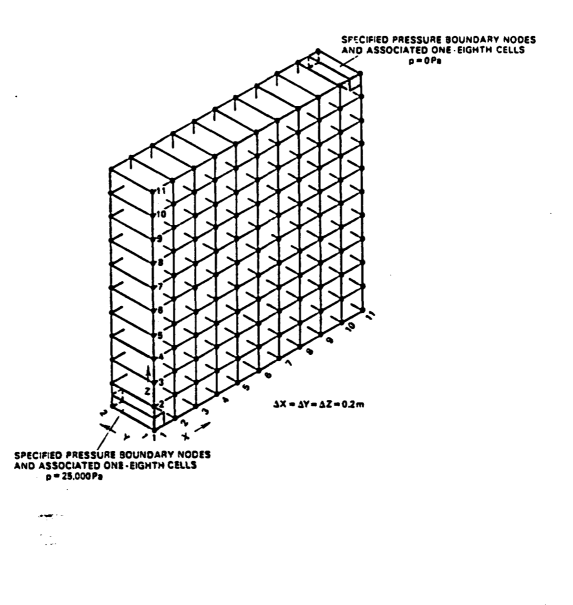


Figure 8.1.--Sketch of the grid with boundary conditions for example problem 1.

Table 8.5.--Input-data file for example problem 1

C.....HST DATA-INPUT FORM C... NOTES: C...: INPUT LINES ARE DENOTED BY C.N1.N2.N3 WHERE NI IS THE READ GROUP NUMBER, NZ.N3 IS THE RECORD NUMBER C... A LETTER INDICATES AN EXCLUSIVE RECORD SELECTION MUST BE MADE C... C... I.E. A OR B OR C (0) - OPTIONAL DATA WITH CONDITIONS FOR REQUIREMENT C... A RECORD NUMBER IN SQUARE BRACKETS IS THE RECORD WHERE THAT PARAMETER IS FIRST SET °C... C..... INPUT BY I.J.K RANGE FORMAT IS: C.O.1.. 11.12.J1.J2.K1.K2 C.O.2.. VAR1, [HOD1, [VAR2, [HOD2, VAR3, [HOD3] C... USE AS MANY OF LINE 0.1 & 0.2 SETS AS NECESSARY C... END WITH LINE 0.3 C.O.3.. 0 / THE SPACE IS REQUIRED C... (NNN) - INDICATES THAT THE DEFAULT NUMBER, NNN, IS USED C... IF A ZERO IS ENTERED FOR THAT VARIABLE (T/F) - INDICATES A LOGICAL VARIABLE C... C... [1] - INDICATES AN INTEGER VARIABLE C....START OF THE DATA FILE C....DIMENSIONING DATA - READI C.1.1 .. TITLE LINE 1 EXAMPLE #1 SOLUTE TRANSPORT WITH VARIABLE DENSITY AND VISCOSITY C.1.2 .. TITLE LINE 2 DIAGONAL FLOW IN X-Z PLANE C.1.3 .. RESTRT(T/F), TIMRST F / C.1.4 .. HEAT, SOLUTE, EEUNIT, CYLIND, SCALMF ; ALL (T/F) FTFFT C.1.5 .. NX,NY,NZ,MHCN 11 2 11 0 C.1.6 .. NPTCBC, NFBC, NAIFC, NLBC, NHCBC, NVEL 4 5\*0 C.1.7 .. NPHZ C.1.8 .. SLMETH[I],LCROSD(T/F) 1 T C.1.9 .. IBC BY I, J, K RANGE (0.1-0.3) , WITH NO IMOD PARAMETER , FOR EXCLUDED CELLS 0/ C.1.10 .. RDECHO(T/F) C....STATIC DATA - READ2 C....OUTPUT INFORMATION C.2.1 .. PRTRE(T/F)

Table 8.5.--Input-data file for example problem 1--Continued

.

```
F
C....COORDINATE GEONETRY INFORMATION
C..... RECTANGULAR COORDINATES
C.2.2A.1 .. UNIGRX_UNIGRY_UNIGRZ: ALL (T/F) ; (0) - NOT CYLIND [1.4]
3*T
C.2.2A.2A .. X(1),X(NX) ;(0) - UNIGRX [2.2A.1]
0. 2.
C.2.2A.2B .. X(1) ;(0) - NOT UNIGRX [2.2A.1]
C.2.2A.3A .. Y(1), Y(NY) ;(0) - UNIGRY [2.2A.1]
0. .2
C.2.2A.38 .. Y(J) ;(0) - NOT UNIGRY [2.2A.1]
C.2.2A.4A .. Z(1), Z(NZ) ;(0) - UNIGRZ [2.2A.1]
0. 2.
C.2.2A.4B .. Z(K) ;(0) - NOT UNIGRZ [2.2A.1]
C.2.28.38 .. Z(K) :(0) - NOT UNIGRZ [2.28.3A] CYLIND [1.4]
C.2.3.1 .. TILT(T/F) ;(0) - NOT CYLIND [1.4]
C.2.3.2 .. THETXZ, THETYZ, THETZZ; (0) - TILT [2.3.1] AND NOT CYLIND [1.4]
C....FLUID PROPERTY INFORMATION
C.2.4.1 .. BP
0.
C.2.4.2 .. PRDEN.TRDEN.WO.DENFO
0. 20. 0. 800.
C.2.4.3 .. W1, DENF1 :(0) - SOLUTE [1.4]
.005 1000.
C.2:5.1 .. NOTVO,TVFO(I),VISTFO(I), I=1 TO NOTVO;(0) - HEAT [1.4] OR HEAT [1.4] AND SOLUTE [1.4] OR .NOT.HEAT AND .NOT.SOLUTE [1.4]
C.2.5.2 .. NOTV1, TVF1(1), VISTF1(1), I=1 TO NOTV1; (0) - SOLUTE [1.4] AND HEAT [1.4]
C.2.5.3 .. NOCV, TRVIS, CVIS(I), VISCTR(I) , I=1 TO NOCV; (0) - SOLUTE [1.4]
2 20. 0. .0013 1. .0007
C.....REFERENCE CONDITION INFORMATION
C.2.6.1 .. PAATH
0.
C.2.6.2 .. PO,TO
0. 20.
C.....SOLUTE INFORMATION
C.2.8 .. DH, DECLAH ;(0) - SOLUTE [1.4]
0. 0.
C..... POROUS NEDIA ZONE INFORMATION
C.2.9.1 .. IPWZ, I1Z(IPWZ), 12Z(IPWZ), J1Z(IPWZ), J2Z(IPWZ), K1Z(IPWZ), K2Z(IPWZ)
1 1 11 1 2 1 11
C..... USE AS HANY 2.9.1 LINES AS NECESSARY
C.2.9.2 .. END WITH 0 /
0/
C.....POROUS MEDIA PROPERTY INFORMATION
C.2.10.1 .. KXX(IPHZ), KYY(IPHZ), KZZ(IPHZ), IPHZ=1 TO NPHZ [1.7]
                                                                                 1
```

Table 8.5.--Input-data file for example problem 1--Continued

3\*1.E-8 and a final frame C.2.10.2 .. POROS(IPMZ), IPMZ=1 TO NPMZ [1.7] .1 C.2.10.3 .. ABPH(IPHZ), IPHZ=1 TO NPHZ [1.7] 0. C.....POROUS MEDIA SOLUTE AND THERMAL DISPERSION INFORMATION C.2.12 .. ALPHL(IPHZ), ALPHT(IPHZ), IPHZ=1 TO NPHZ [1.7]; (0) - SOLUTE [1.4] OR HEAT [1.4] .1 .1 C.....POROUS MEDIA SOLUTE PROPERTY INFORMATION C.2.13 .. OBKD(IPHZ) , IPHZ=1 TO MPHZ [1.7] ;(0) - SOLUTE [1.4] 0. C.....BOUNDARY CONDITION INFORMATION C.... SPECIFIED VALUE B.C. C.2.15 .. IBC BY I, J, K RANGE (0.1-0.3) WITH NO IMOD PARAMETER, ;(0) - NPTCBC [1.6] > 0 111211 101 11 11 1 2 11 11 100 0/ C....FREE SURFACE B.C. C.2.20 .. FRESUR(T/F), PRTCCM(T/F) F / \_ C....INITIAL CONDITION INFORMATION C.2.21.1 .. ICHYDP, ICT, ICC; ALL (T/F); IF NOT. HEAT, ICT = F, IF NOT. SOLUTE, ICC = F TFT C.2.21.2 .. ICHWT(T/F) ;(0) - FRESUR [2.20] C.2.21.3A .. ZPINIT PINIT :(0) - ICHYDP [2.21.1] AND NOT ICHWT [2.21.2] 2. 0. C.2.21.6 .. C BY I, J, K RANGE (0.1-0.3) ;(0) - SOLUTE [1.4] AND ICC [2.21.1] 1 11 1 2 1 11 0.1 0/ C....CALCULATION INFORMATION C.2.22.1 .. FDSMTH, FDTMTH .5.5 C.2.22.2 .. TOLDEN(.001), MAXITN(5) 0.0 C....OUTPUT INFORMATION C.2.23.1 .. PRTPMP, PRTFP, PRTIC, PRTBC, PRTSLM, PRTWEL; ALL (T/F) . . . . . . . C.2.23.2 .. IPRPIC, PRIDV(T/F) ;(0) - PRIIC [2.23.1] 101 T C.2.23.3 .. ORENPR[1]; (0) - NOT CYLIND [1.4] 13 C.2.23.4 .. PLTZON(T/F)

C.2.23.5 .. OCPLOT(T/F) S. à. C-----\_\_\_\_\_ C..... TRANSIENT DATA - READ3 C.3.1 .. THRU(T/F) C....IF THRU IS TRUE PROCEED TO RECORD 3.99 C.....THE FOLLOWING IS FOR NOT THRU C....BOUNDARY CONDITION INFORMATION SPECIFIED VALUE B.C. C.... C.3.3.1 .. RDSPBC, RDSTBC, RDSCBC, ALL(T/F); (0) - NOT CYLIND [1.4] AND NPTCBC [1.6] > 0 OR CYLIND AND NPTCBC > 1 TFT C.3.3.2 .. PNP B.C. BY I.J.K RANGE (0.1-0.3) ;(0) - RDSPBC [3.3.1] 111211 25000.1 11 11 1 2 11 11 0.1 0/ C.3.3.4 .. CSBC BY I, J, K RANGE (0.1-0.3); (0) - RDSPBC [3.3.1] AND SOLUTE [1.4] 0/ C.3.3.6 .. CNP B.C. BY I.J.K RANGE (0.1-0.3) :(0) - RDSCBC [3.3.1] AND SOLUTE [1.4] 111211 1.1 0/ C....CALCULATION INFORMATION C.3.7.1 .. RDCALC(T/F) C.3.7.2 .. AUTOTS(T/F) ;(0) - ROCALC [3.7.1] C.3.7.3.A .. DELTIN :(0) - RDCALC [3.7.1] AND NOT AUTOTS [3.7.2] .2 C.3.7.4 .. TINCHG 10. C....OUTPUT INFORMATION C.3.8.1 .. PRIVEL, PRIDV, PRISLM, PRIKD, PRIPTC, PRIGFB, PRIWEL, PRIBCF ; ALL [1] 8\*-1 C.3.8.2 .. IPRPTC :(0) - IF PRIPTC [3.8.1] NOT = 0 101 C.3.8.3 .. CHKPTD(T/F), HTSCHK, SAVLDO(T/F) F / C....CONTOUR MAP INFORMATION C.3.9.1 .. RDMPDT, PRTMPD; ALL (T/F) TT C.3.9.2 .. NAPPTC, PRIMAP[1] :(0) - RDMPDT [3.9.1]

.

Table 8.5.--Input-data file for example problem 1--Continued

Table 8.5.--Input-data file for example problem 1--Continued

1.4 001 -1 C.3.9.3 .. YPOSUP(T/F), ZPOSUP(T/F), LENAX, LENAY, LENAZ; (0) - ROMPDT [3.9.1] FT 10. 0. 10. C.3.9.4 .. IMAP1(1), IMAP2(NX), JMAP1(1), JMAP2(NY), KMAP1(1), KMAP2(NZ), AMIN, AMAX, NMPZON(5):(0) - RDMPDT [3.9.1] 1 11 1 1 1 11 0. 1. 10 C....ONE OF THE 3.9.4 LINES REQUIRED FOR EACH DEPENDENT VARIABLE C..... TO BE MAPPED C....END OF FIRST SET OF TRANSIENT INFORMATION C.....READ SETS OF READ3 DATA AT EACH TINCHG UNTIL THRU (LINES 3.N1.N2) C....END OF CALCULATION LINES FOLLOW, THRU-.TRUE. . . . C.3.99.1 .. THRU T. C....TEMPORAL PLOT INFORMATION C.3.99.2 .. PLOTWP, PLOTWT, PLOTWC; ALL (T/F) 3\*F

C....END OF DATA FILE

Table 8.6.--Output file for example problem 1

THREE DIMENSIONAL FLOW, HEAT AND SOLUTE TRANSPORT SIMULATOR - (HST3D):RELEASE - 1.0

#### EXAMPLE #1 SOLUTE TRANSPORT WITH VARIABLE DENSITY AND VISCOSITY DIAGONAL FLOW IN X-Z PLANE

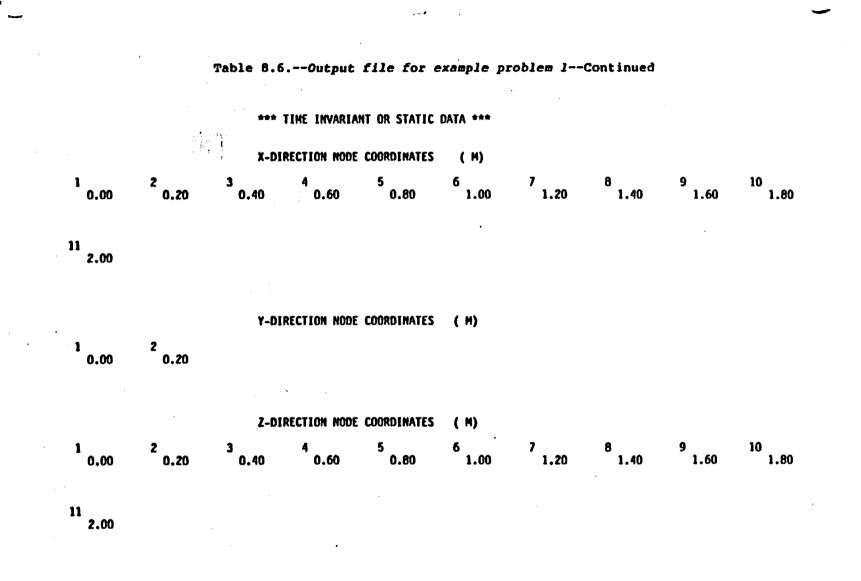
\*\*\* FUNDAMENTAL INFORMATION \*\*\* CARTESIAN COORDINATES ISOTHERMAL SIMULATION SOLUTE TRANSPORT SIMULATION INPUT DATA IS EXPECTED IN METRIC UNITS SOLUTE CONCENTRATION IS EXPRESSED AS SCALED MASS FRACTION WITH RANGE (0-1)

\*\*\* PROBLEM DIMENSIONING INFORMATION \*\*\*

NUMBER	QF	NODES IN X-DIRECTION	NX	11
NUMBER	OF	NODES IN Y-DIRECTION	NY	2
		NODES IN Z-DIRECTION		11
NUMBER	OF	POROUS MEDIA ZONES	NPHZ .	1
NUMBER	OF	SPECIFIED PRESSURE, TEMPERATURE OR MASS FRACTION B.C.	NPTCBC	4
NUMBER	ŌF	SPECIFIED FLUX B.C. CELLS (FLOW, HEAT OR SOLUTE)	NFBC .	0
		HEAT CONDUCTION B.C. CELLS		0
NUMBER	QE.	NODES OUTSIDE REGION FOR EACH HEAT CONDUCTION B.C. CELL	NHCN	0
NUMBER	ÔF	AQUIFER INFLUENCE FUNCTION CELLS	NALFC	0
		LEAKAGE CELLS		Ó
		WELLS		Ō

### DIRECT D4 SOLVER IS SELECTED

ABBREVIATED DIAGONAL CROSS-DISPERSIVITY COEFFICIENT STORAGE ALLOC	ATED	
THE A4 ARRAY IN DADES IS DIMENSIONED		3799 ELEMENTS
THE TOTAL STORAGE REQUIRED BY THE DIRECT METHOD IS		5009 ELEMENTS
THE TOTAL STORAGE REQUIRED BY THE ITERATIVE METHOD IS		1694 ELEMENTS
TOTAL LENGTH OF LABELED COMMON BLOCKS		6681 BYTES
	REQUIRED	COMPILED
LENGTH OF VARIABLE LENGTH REAL ARRAY (VPA ARRAY) 13	1076 ELEMENTS	250000 ELEMENTS
LENGTH OF VARIABLE LENGTH INTEGER ARRAY (IVPA ARRAY) 1	779 ELEMENTS	20000 ELEMENTS



<u>\*</u> -



REGION 11 12 J1 J2 K1	POROUS MEDIUM K2 ZONE INDEX	•
1 11 1 2 1		•••••••••••••••••••••••••••••••••••••••

**\*\*\* POROUS NEDIA PROPERTIES \*\*\*** 

X-DIRECTION PERHEABILITIES ( N\*\*2)

1 1.0000E-08

Y-DIRECTION PERHEABILITIES ( M\*\*2)

1 1.0000E-08

Z-DIRECTION PERMEABILITIES ( H\*\*2)

1 1.0000E-08

POROSITY (-)

1 0.1000

1

\*\*\* INTERMEDIATE COMPUTED DATA \*\*\*

4

X-DIRECTION CONDUCTANCE FACTOR BETWEEN X(I) AND X(I+1) ( N\*\*3)

VERTICAL SLICES

3

2

J = 1

6

5

7 8

11 5.0000E-10 10 1.0000E-09 9 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1,0000E-09 8 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 7 1.0000E-09 6 1.00008-09 1.0000E-09 1.0000£-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 2 1.0000E-09 

J = 2

	1	. <b>2</b>	3	4	5	6	7	8	9	10
11	5.0000E-10	5,0000E-10	5.00008-10							
10	1.0000E-09	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09
9	1.0000E-09									
8	1.0000E-09									
7	1.0000E-09	1.0000E-09	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09	1.0000E-09
. 6	1.0000E-09									
5.	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09						
4	1.0000E-09									
. 3	1.0000E-09									
2	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.00008-09	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09
1	5.0000E-10									

Y-DIRECTION CONDUCTANCE FACTOR BETWEEN Y(J) AND Y(J+1) ( M\*\*3)

VERTICAL SLICES

J = 1

2 5 6 A 10 11 5.0000E-10 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 1.0000E-09 10 1.0000E-09 2.0000E-09 9 1.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 8 1.0000E-09 2.0000E-09 7 1.0000E-09 2.0000E-09 2.00008-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 6 1.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.00008-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 5 1.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 4 1.0000E-09 2.0000E-09 3 1.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09 2.0000E-09

.

2 1	1.0000E-09 5.0000E-10	2.0000E-09 1.0000E-09									
11 10 9	11 5.0000E-10 1.0000E-09 1.0000E-09										
8	1.0000E-09										
7	1.0000E-09										
6	1.0000E-09										
- 5	1.00008-09								•		
4	1.00008-09										
3	1.0000E-09										
•	1 00006 00										

2 1.0000E-09 1 5.0000E-10

Z-DIRECTION CONDUCTANCE FACTOR BETWEEN Z(K) AND Z(K+1) ( H\*\*3)

VERTICAL SLICES

J = 1

	1	2	3	. 4	5	6	7	8	9	10
10	5.0000E-10	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.00008-09	1.00008-09
9	5.0000E-10	1.0000E-09	1.00002-09							
8	5.0000E-10	1.0000E-09								
7	5.0000E-10	1.0000E-09								
6	5.0000E-10	1.0000E-09								
5	5.0000E-10	1.00008-09	1.0000E-09	1.0000E-09	1.00002-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09
4	5.0000E-10	1.0000E-09								
3	5.0000E-10	1.00008-09	1.0000E-09							
2	5.0000E-10	1.0000E-09								
1	5.0000E-10	1.0000E-09								

11 10 5.0000E-10 9 5.0000E-10 8 5.0000E-10 7 5.0000E-10

. .

6 5.0000E-10 5 5.0000E-10 4 5.0000E-10 3 5.0000E-10 2 5.0000E-10 1 5.0000E-10

J = 2

.

	1	2	3	4 .	5	6	7	8	9	10
10	5.0000E-10	1.0000E-09	1.00008-09	1.0000E-09						
9	5.0000E-10	1.0000E-09								
8	5.0000E-10	1.0000E-09								
7	5.0000E-10	1.0000E-09	1,0000E-09	1.0000E-09						
6	5.0000E-10	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.00008-09	1.0000E-09	1.0000E-09
5	5.0000E-10	1.0000E-09	1.0000E-09	1.0000E-09	1,0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09
- 4	5.0000E-10	1.0000E-09	1,0000E-09	1.0000E-09	1.0000E-09	1,0000E-09	1.0000E-09	1.0000E-09	1.0000E-09	1.0000E-09
3	5.0000E-10	1.0000E-09								
2	5.0000E-10	1.0000E-09								
1	5.0000E-10	1.0000E-09								

261

11 10 5.0000E-10 5.0000E-10 9 5.0000E-10 8 5,0000E-10 7 5.0000E-10 6 5.0000E-10 5 5.0000E-10 4 5.0000E-10 3 2 5.0000E-10 1 5.0000E-10

#### \*\*\* PROPERTIES BY POROUS MEDIUM ZONE \*\*\*

ł

# POROUS MEDIUM VERTICAL COMPRESSIBILITY (1/ PA)

1 0.0000

LONGITUDINAL DISPERSIVITY ( H)

1 0.1000

TRANSVERSE DISPERSIVITY ( M)

.

1 0.1000

DENSITY - DISTRIBUTION COEFFICIENT PRODUCT (-)

1 0.0000

 NOLECULAR DIFFUSIVITY-TORTUOSITY PRODUCT
 DM
 0.000
 (M\*\*2/S)

 SOLUTE LINEAR DECAY RATE CONSTANT
 DECLAH 0.000
 (1/S)

 SCALE FACTORS FOR SCALED MASS FRACTION
 NO
 0.00000

 N1
 0.00500

 ATHOSPHERIC PRESSURE (ABSOLUTE)
 PAATH
 101325.0
 (PA)

 REFERENCE PRESSURE FOR ENTHALPY
 POH
 0.0
 (PA)

 ISOTHERNAL AQUIFER TEMPERATURE
 TOH
 20.0
 (DEG.C)

VISCOSITY-CONCENTRATION DATA TABLE AT 20.0 DEG.C SCALED MASS FRACTION VISCOS

VISCOSITY (KG/ M-S)

.

0.00	1.300E-03
1.00	7.0008-04

\*\*\* INITIAL CONDITIONS \*\*\*

INITIAL AQUIFER FLUID PRESSURE FOR HYDROSTATIC I.C PINIT.	· 0.0 (	( PA)
ELEVATION OF INITIAL PRESSURE	2.0	(H)

INITIAL PRESSURE DISTRIBUTION ( PA)

.

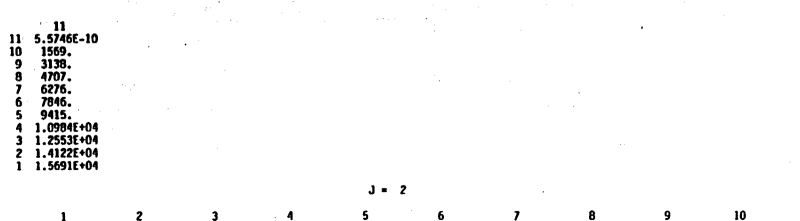
VERTICAL SLICES

÷ .

,

J = 1

	1	2	3	4	5	6	7	8	9	10
11	5.5746E-10									
10	1569.	1569.	1569.	1569.	1569.	1569.	1569.	1569.	1569.	1569.
9	3138.	3138.	3138.	3138.	3138.	3138.	3138.	3138.	3138.	3138.
8	4707.	4707.	4707.	4707.	4707.	4707.	4707.	4707.	4707.	4707.
7	6276.	6276.	6276.	6276.	6276.	6276.	6276.	6276.	6276.	6276.
6	7846.	7846.	7846.	7846.	7846.	7846.	7846.	7846.	7846.	7846.
5	9415.	9415.	9415.	9415.	9415.	9415.	9415.	9415.	9415.	9415.
- 4	1.0984E+04									
3	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04		1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04
2	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04		1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04
1	1.5691E+04									



2 3 5 7 8 9 1 . 4 6

263

.

11	5.5746E-10	5.5746E-10	5.5746E-10	5.57468-10	5.5746E-10	5.57462-10	5.5746E-10	5.5746E-10	5.5746E-10	5.5746E-10
10	1569.	1569.	1569.	1569.	1569.	1569.	1569.	1569.	1569.	1569.
9	3138.	3138.	3138.	3138.	3138.	3138.	3138.	3138.	3138.	3138.
8	4707.	4707.	4707.	4707.	4707.	4707.	4707.	4707.	4707.	4707.
7	6276.	6276.	6276.	6276.	6276.	6276.	6276.	6276.	6276.	6276.
6	7846.	7846.	7846.	7846.	7846.	7846.	7846.	7846.	7846.	7846.
5	9415.	<del>9</del> 415.	9415.	9415.	9415.	9415.	9415.	9415.	9415.	9415.
- 4	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04	1.0984E+04
3	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04	1.2553E+04
2	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04	1.4122E+04
1	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04	1.5691E+04

11 11 5.5746E-10 10 1569. 3138. 4707. 9 8 6276. 7 7846. 6 9415. 1.0984E+04 4

1.2553E+04

2 1.4122E+04 1 1.5691E+04

# INITIAL SCALED HASS FRACTIONS ( - )

VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
11	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
10	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
9	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
8	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
7	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
6	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
5	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
4	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0,0000
3	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000

2	0.0000 0.0000	0.0000	0.0000	0.0000 0.0000	0.0000 0.0000	0.0000 0.0000	0.0000 0.0000	0.0000 0.0000	0.0000 0.0000	0.0000 0.0000
•		•								
11 10 9 8 7 6 5 4 3 2	11 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	·								·
1	0.0000	· · ·	. 4	1						
			. • * *		J = 2					
11 10 9 7 6 5 4 3 2 1	1 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	2 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	3 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	4 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	5 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	6 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	7 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	8 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	9 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	10 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000

 $\begin{array}{c|ccccc} & 11 \\ 11 & 0.0000 \\ 10 & 0.0000 \\ 9 & 0.0000 \\ 8 & 0.0000 \\ 7 & 0.0000 \\ 6 & 0.0000 \\ 6 & 0.0000 \\ 5 & 0.0000 \\ 4 & 0.0000 \\ 3 & 0.0000 \end{array}$ 

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. .

2 0.0000 1 0.0000

### INITIAL PORE VOLUME PER CELL ( H\*\*3)

### VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	.9	10
11	1.0000E-04	2.0000E-04								
10	2.0000E-04	4.0000E-04								
9	2.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.00008-04	4.0000E-04	4.0000E-04	4.0000E-04
8	2.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.00008-04	4.0000E-04	4.0000E-04	4.0000E-04
7	2.0000E-04	4.0000E-04								
6	2.0000E-04	4.0000E-04								
5	2.0000E-04	4.0000E-04								
4	2.0000E-04	4.0000E-04	4,00000-04	4.0000E-04						
3	2.0000E-04	4.0000E-04	4.00008-04	4.0000E-04						
2	2.0000E-04	4.0000E-04								
1	1.0000E-04	2.0000E-04	2.0000E-04	2.00008-04	2.0000E-04	2.0000E-04	2.0000E-04	2.0000E-04	2,0000E-04	2.0000E-04

11 11 1.0000E-04 10 2.0000E-04 9 2.0000E-04 8 2.0000E-04 7 2.0000E-04 6 2.0000E-04 5 2.0000E-04 4 2.0000E-04 3 2.0000E-04

2 2.0000E-04

1 1.00D0E-04

J = 2

 1
 2
 3
 4
 5
 6
 7
 8
 9
 10

 11
 1.0000E-04
 2.0000E-04
 4.0000E-04
 4

ية. ب

							1. A.		2.00	
1	1.0000E-04	2.00002-04	2.0000E-04	2.0000E-04	2.0000E-04	2.00008-04	2.0000E-04	2.00008-04	2.0000E-04	2.0000E-04
						4.0000E-04				
						4.0000E-04				
4	2.0000E-04	4.0000E-04	4.00008-04	4.0000E-04	4,0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04
5	2.0000E-04	4.0000E-04	4.00008-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.00008-04	4.0000E-04	4.0000E-04
6	2.0000E-04	4.00008-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.00002-04	4.0000E-04	4.0000E-04	4.0000E-04
7	2.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04
8	2.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04	4.0000E-04

	11	
11	1.0000E-04	· · · ·
10	2.0000E-04	۰,
. 9	2.0000E-04	
8	2.0000E-04	
7	2.0000E-04	
6	2.0000E-04	
5	2.0000E-04	
4	2.0000E-04	
3	2.0000E-04	
2	2.0000E-04	
1	1.0000E-04	

INITIAL FLUID DENSITY IN CELL (KG/ H\*\*3)

VERTICAL SLICES

A 14

J = 1

	1	2	3	4	5	6	7	8	9	10
- 11	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
10	800,00	800.00	800.00	800.00	.800.00	800.00	800.00	800.00	800.00	800.00
9	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
8	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
7	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
. 6	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
5	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
- 4	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
3	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00
2	800.00	800.00	800.00	800.00	800.00	800.00	800,00	800.00	800.00	800.00
1	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00	800.00

11 10 9 8 7 6 5 4 3 2 1	11 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00											
	J = 2											
11 10 9 8 7 6 5 4 3 2 1	1 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	2 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	3 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	4 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	5 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	6 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	7 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	8 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	9 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00 800.00	$\begin{array}{c} 10\\ 800.00\\ 800$		

.

11 11 800.00 9 800.00 9 800.00 9 800.00 7 800.00 6 800.00 5 800.00 5 800.00 4 800.00 3 800.00 2 800.00 1 800.00

# INITIAL FLUID VISCOSITY IN CELL (KG/ M-S)

VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
11	1.30002+03	1.3000E-03								
10	1.3000E-03									
9	1.3000E-03									
8	1.3000E-03									
7	1.3000E-03									
6	1.3000E-03									
5	1.3000E-03									
4	1.3000E-03									
3	1.3000E-03									
2	1.3000E-03									
1	1.3000E-03									

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.....

11 11 1.3000E-03 10 1.3000E-03 9 1.3000E-03 8 1.3000E-03 7 1.3000E-03 6 1.3000E-03 5 1.3000E-03 4 1.3000E-03 3 1.3000E-03 2 1.3000E-03

1 1.3000E-03

J = 2

	1	2	3	4	5	6	7	8	9	10
11	1.3000E-03									
			1.3000E-03							
9	1.3000E-03									
8	1.3000E-03									
			1.3000E-03							
6	1.3000E-03									

..... 6.400000E+01 (KG) ; 8.000000E-02 ( M\*\*3)

5	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.30008-03	1.30008-03
4	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.30008-03	1.3000E-03
3	1.3000E-03	1. JUUOE-03	1.3000E-03							
					1.3000E-03					
1	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.3000E-03	1.30008-03

J = 1

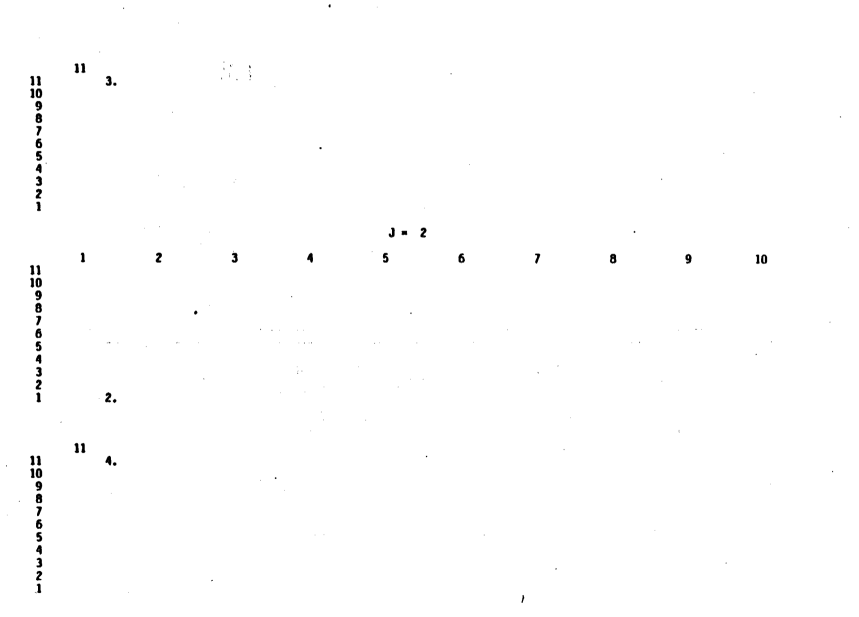
INDEX NUMBERS FOR SPECIFIED P.T OR C NODES

11 1.3000E-03 10 1.3000E-03 9 1.3000E-03 8 1.3000E-03 7 1.3000E-03 6 1.3000E-03 5 1.3000E-03 4 1.3000E-03 4 1.3000E-03 3 1.3000E-03 2 1.3000E-03 1 1.3000E-03

1.

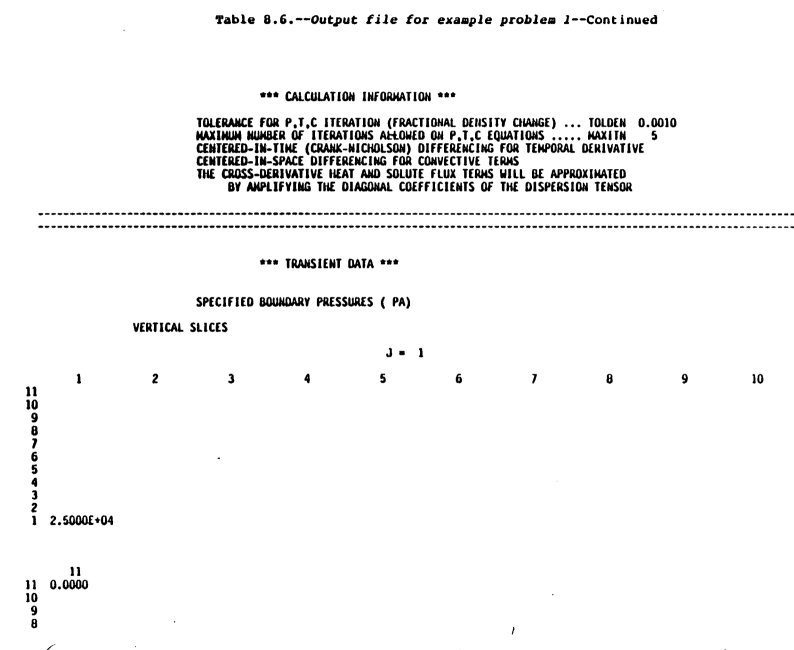
INITIAL FLUID IN REGION ....

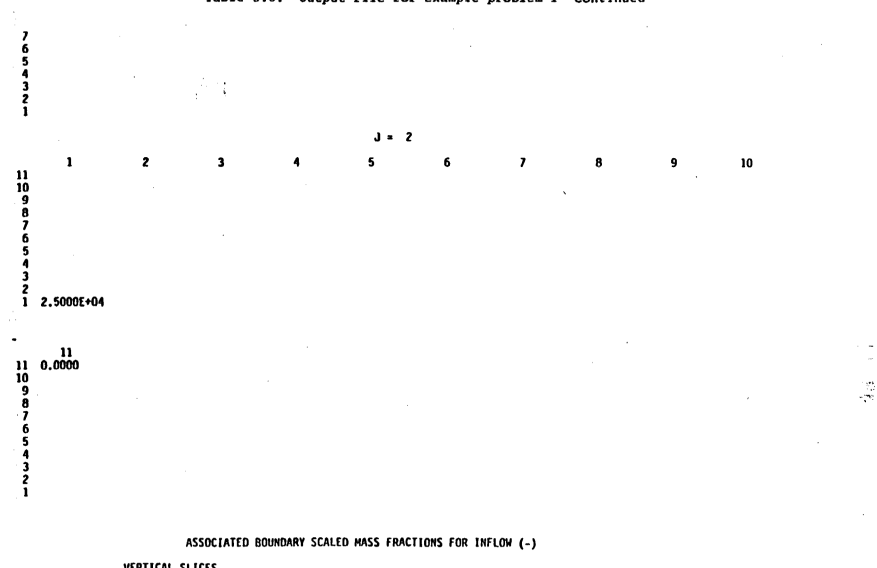
VERTICAL SLICES



....

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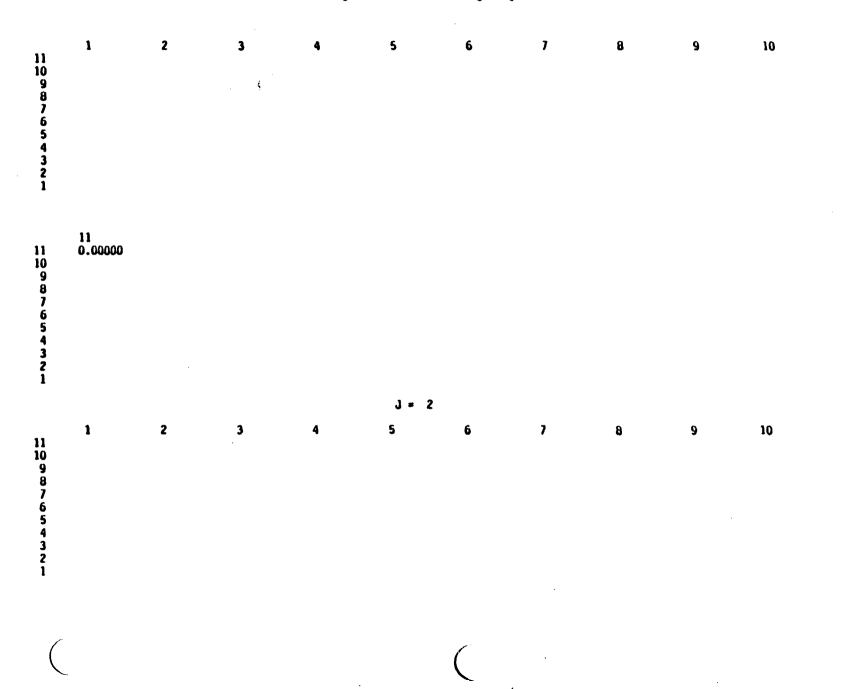




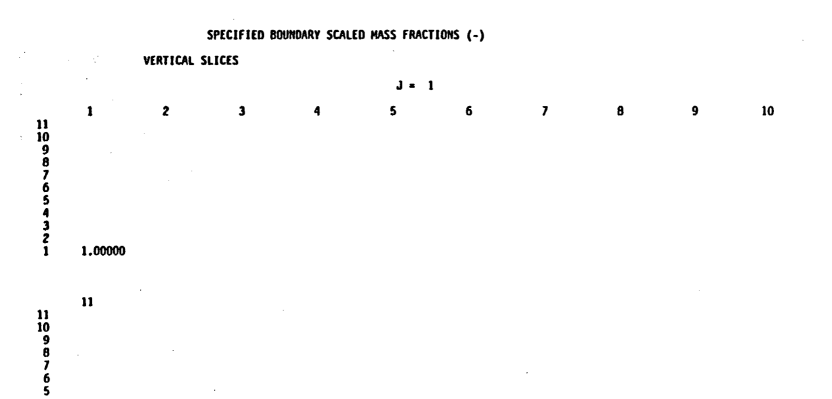
273

VERTICAL SLICES

J = 1







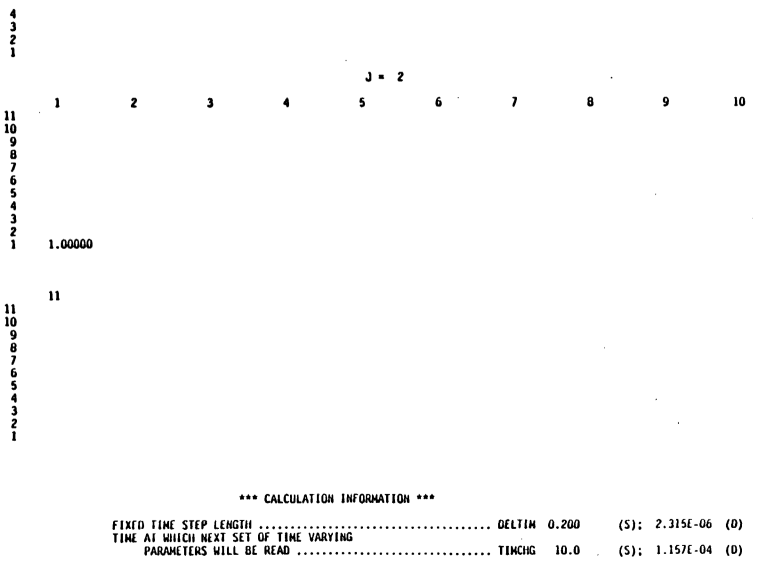


Table 8.6.--Output file for example problem 1--Continued

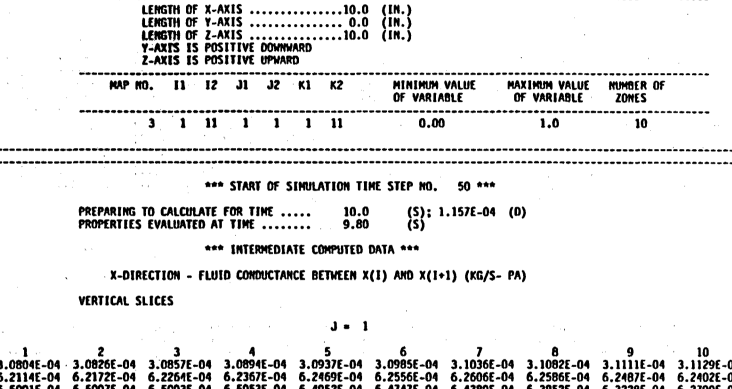
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\*\*\* MAPPING DATA \*\*\*

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 11
 3.0804E-04
 3.0857E-04
 3.0894E-04
 3.0937E-04
 3.0985E-04
 3.1036E-04
 3.1082E-04
 3.111E-04
 3.1129E-04

 10
 6.2172E-04
 6.2264E-04
 6.2367E-04
 6.2469E-04
 6.2556E-04
 6.2606E-04
 6.2586E-04
 6.2487E-04
 6.2402E-04

 9
 6.5091E-04
 6.5097E-04
 6.5093E-04
 6.5053E-04
 6.4747E-04
 6.4389E-04
 6.32586E-04
 6.2487E-04
 6.2402E-04

 8
 7.4652E-04
 7.4200E-04
 7.3384E-04
 7.2310E-04
 7.1038E-04
 6.9561E-04
 6.7854E-04
 6.6088E-04
 6.3228E-04
 6.3308E-04

 7
 9.5025E-04
 9.3328E-04
 9.0359E-04
 8.6606E-04
 8.2429E-04
 7.8020E-04
 7.3532E-04
 6.9304E-04
 6.5929E-04
 6.4045E-04

 6
 1.2159E-03
 1.1878E-03
 1.0703E-03
 9.9128E-04
 9.0534E-04
 8.1901E-04
 7.4129E-04
 6.8256E-04
 6.5118E-04

 5
 1.3823E-03
 1.3229E-03
 1.2602E-03
 1.1718E-03
 1.0584E-03
 9.3039E-04
 8.0881E-04
 7.1586E-04
 6.6631E-04

 4
 1.4234E-03
 1.4101E-03
 1.

J = 2

1 2 3 4 5 6 7 8 9 10 11 3.0804E-04 3.0826E-04 3.0857E-04 3.0894E-04 3.0937E-04 3.0985E-04 3.1036E-04 3.1082E-04 3.1111E-04 3.1129E-04

1 7.1428E-04 7.1421E-04 7.1361E-04 7.1064E-04 7.0108E-04 6.7855E-04 6.3621E-04 5.6935E-04 4.8063E-04 3.9647E-04

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.

10	6.2114E-04	6.2172E-04	6.2264E-04	6.2367E-04	6.2469E-04	6.2556E-04	6.2606E-04	6.2586E-04	6.2487E-04	6.2402E-04
9	6.5091E-04	6.5097E-04	6.5093E-04	6.5053E-04	6.4952E-04	6.4747E-04	6.4389E-04	6.3852E-04	6.3228E-04	6.2790E-04
8	7.4652E-04	7.4200E-04	7.3384E-04	7.2310E-04	7.10388-04	6.9561E-04	6.7854E-04	6.6008E-04	6.4332E-04	6.3308E-04
7	9.5025E-04	9.3328E-04	9.0359E-04	8.6606E-04	8.2429E-04	7.8020E-04	7.3532E-04	6.9304E-04	6.5929E-04	6.4045E-04
6	1.2159E-03	1.18782-03	1.1374E-03	1.0703E-03	9.9128E-04	9.0534E-04	8.1901E-04	7.4129E-04	6.8256E-04	6.5118E-04
5	1.3823E-03	1.3628E-03	1.3229E-03	1.2602E-03	1.1718E-03	1.0584E-03	9.3039E-04	8.0881E-04	7.1586E-04	6.6631E-04
4	1.4234E-03	1.4171E-03	1.4001E-03	1.3648E-03	1.3018E-03	1.2004E-03	1.0580E-03	8.9748E-04	7.6207E-04	6.8671E-04
3	1.4281E-03	1.4267E-03	1.4212E-03	1.4050E-03	1.3675E-03	1.2950E-03	1.1729E-03	1.0010E-03	8.2402E-04	7.1332E-04
		1.4282E-03					·1.2464E-03			
1	7.1428E-04	7.1421E-04	7.1361E-04	7.1064E-04	7.0108E-04	6.7855E-04	6.3621E-04	5.6935E-04	4.8063E-04	3.9647E-04

Y-DIRECTION - FLUID CONDUCTANCE BETWEEN Y(J) AND Y(J+1) (KG/S- PA)

VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
11	3.0796E-04	6.1625E-04	6.1680E-04	6.1749E-04	6.1829E-04	6.1921E-04	6.2021E-04	6.2123E-04	6.2203E-04	6.2242E-04
10	6.2098E-04	1.2426E-03	1.2443E-03	1.2463E-03	1.2484E-03	1.2504E-03	1.2519E-03	1.2523E-03	1.2511E-03	1.2484E-03
9	6.5089E-04	1.30198-03	1.3020E-03	1.3017E-03	1.3004E-03	1.2976E-03	1.2923E-03	1.2833E-03	1.2708E-03	1.2584E-03
8	7.4773E-04	1.4906E-03	1.4774E-03	1.4581E-03	1.4345E-03	1.4072E-03	1.3755E-03	1.3391E-03	1.30162-03	1.2719E-03
7	9.54898-04	1.8913E-03	1.8423E-03	1.7730E-03	1.6926E-03	1.6062E-03	1.5164E-03	1.4268E-03	1.3470E-03	1.2910E-03
6	1.2236E-03	2.4166E-03	2.3356E-03	2.2163E-03	2.0687E-03	1.9016E-03	1.72COE-03	1.5565E-03	1.4137E-03	1.31 <del>89</del> E-03
5	1.3873E-03	2.7548E-03	2.6967E-03	2.5962E-03	2.4477E-03	2.2461E-03	1.9982E-03	1.7368E-03	1.5100E-03	1.3591E-03
4	1.4247E-03	2.8442E-03	2.8243E-03	2.7765E-03	2.6839E-03	2.5267E-03	2.2842E-03	1.9656E-03	1.6452E-03	1.4156E-03
3	1.4283E-03	2.8558E-03	2.85128-03	2.8339E-03	2.78652-03	2.6848E-03	2.5000E-03	2.2053E-03	1.8253E-03	1.4948E-03
2	1.4285E-03	2.8570E-03	2.85598-03	2.8492E-03	2.82388-03	2.75608-03	2.61668-03	2.3774E-03	2.0233E-03	1.6099E-03
1	7.1429E-04	1.4285E-03	1.4283E-03	1.4261E-03	1.4165E-03	1.3881E-03	1.3271E-03	1.2210E-03	1.0644E-03	8.7219E-04

11 11 3.1137E-04 10 6.2382E-04 9 6.2662E-04 8 6.3023E-04 7 6.3544E-04 6 6.4303E-04 5 6.5342E-04 4 6.6651E-04 3 6.8144E-04 2 727E-04

1 3.6187E-04

VERTICAL SLICES

 $\{\lambda_i\}_{i=1}^{n}$ 

				<b>J</b> = 1				i.	•	;
	1	2	3	4	5	6	7	8	9	10
10	3.0922E-04						6.2307E-04		6.2378E-04	
9	3.1784E-04	6.3587E-04	6.3634E-04	6.3678E-04	6.3702E-04	6.3684E-04	6.3593E-04	6.3385E-04	6.3044E-04	6.2669E-04
8	3.4845E-04	6.9582E-04	6.9287E-04	6.8835E-04	6.8255E-04	6.7542E-04	6.6647E-04	6.5538E-04	6.4305E-04	6.3255E-04
7	4.2104E-04	8.3681E-04	8.2261E-04	8.0218E-04	7.7789E-04	7.5099E-04	7.2173E-04	6.9097E-04	6.6202E-04	6.4070E-04
6	5.3846E-04	1.0651E-03	1.03372-03	9.8823E-04	9.3338E-04	8.7242E-04	8.0813E-04	7.44808-04	6.8987E-04	6.5243E-04
5	6.5080E-04	1.2887E-03	1.2532E-03	1.1975E-03	1.1232E-03	1.0316E-03	9.2739E-04	8.2151E-04	7.3034E-04	6.6941E-04
4	7.02898-04	1.3995E-03	1.3797E-03	1.3420E-03	1.2809E-03	1.1901E-03	1.0671E-03	9.2300E-04	7.8775E-04	6.9348E-04
3	7.1324E-04	1.4250E-03	1.4188E-03	1.4025E-03	1.3672E-03	1.3020E-03	1.1942E-03	1.04028-03	8.6591E-04	7.2720E-04
2	7.1421E-04	1.4282E-03	1.4268E-03	1.4208E-03	1.4025E-03	1.3600E-03	1.2786E-03	1.1445E-03	9.6028E-04	7.7539E-04
÷ 1.	7.1428E-04	1.4285E-03	1.4281E-03	1.4254E-03	1.4142E-03	1.3830E-03	1.3177E-03	1.2047E-03	1.0375E-03	8.3758E-04

	1. A 11. A
	11 1
10	3.1164E-04
9	3.12618-04
8	3.1421E-04
7	3.1641E-04
6	3.1961E-04
5	3.2410E-04
Ā	3.2996E-04
3	3.36962-04
2	3.4465E-04
៍រិ	3.5517E-04

J = 2

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 10
 3.0922E-04
 6.1877E-04
 6.1946E-04
 6.2031E-04
 6.2123E-04
 6.2219E-04
 6.2307E-04
 6.2369E-04
 6.2378E-04
 6.2332E-04

 9
 3.1784E-04
 6.3587E-04
 6.3634E-04
 6.3678E-04
 6.3702E-04
 6.3684E-04
 6.3593E-04
 6.3385E-04
 6.2369E-04
 6.3647E-04
 6.3593E-04
 6.3044E-04
 6.2669E-04

 8
 3.4845E-04
 6.95882E-04
 6.8835E-04
 6.8255E-04
 6.7542E-04
 6.6647E-04
 6.5538E-04
 6.4305E-04
 6.3255E-04

 7
 4.2104E-04
 8.3681E-04
 8.0218E-04
 7.7789E-04
 7.5099E-04
 7.2173E-04
 6.9097E-04
 6.6202E-04
 6.4070E-04
 6.5243E-04
 6.5243E-04
 5.5886E-

4	7.0289E-04	1.39958-03	1.37978-03	1.3420E-03	1.2809E-03	1.1901E-03	1.0671E-03	9.2300E-04	7.8775E-04	6.9348E-04
3	7.1324E-04	1.4250E-03	1.4188E-03	1.4025E-03	1.3672E-03	1.3020E-03	1.1942E-03	1.0402E-03	8.6591E-04	7.2720E-04
2	7.1421E-04	1.4282E-03	1.4268E-03	1.4208E-03	1.4025E-03	1.3600E-03	1.2786E-03	1.1445E-03	9,6028E-04	7.7539E-04
1	7.1428E-04	1.4285E-03	1.4281E-03	1.4254E-03	1.4142E-03	1.3830E-03	1.3177E-03	1.2047E-03	1.0375E-03	8.3/58E-04

11 10 3.1164E-04 9 3.1261E-04 8 3.1421E-04 7 3.1641E-04 6 3.1961E-04 5 3.2410E-04 4 3.2996E-04 3 3.3696E-04 2 3.4465E-04 1 3.5517E-04

X-DIRECTION - SOLUTE DISPERSIVE CONDUCTANCE BETWEEN X(I) AND X(I+1) (KG/S)

VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
- 11	2.1266E-03	6.4845E-03	1.1234E-02	1.6811E-02	2.3920E-02	3.3821E-02	4.9145E-02	7.6615E-02	0.1374	0.3114
10	9.3831E-03	1.5388E-02	2.3889E-02	3.4464E-02	4.8098E-02	6.67728-02	9.4014E-02	0.1364	0,2079	0.3609
9	1.7300E-02	2.1333E-02	2.8260E-02	3.7634E-02	4.99868-02	6.6585E-02	8.9262E-02	0.1202	0.1611	0.2091
8	2.67508-02	3.0016E-02	3.5810E-02	4.3767E-02	5.4269E-02	6.8151E-02	8.6094E-02	0.1079	0.1314	0.1508
7	4.0235E-02	4.3271E-02	4.7976E-02	5.37098-02	6.1087E-02	7.1188E-02	8.4360E-02	9.9377E-02	0.1135	0.1230
6	6.1574E-02	6.4471E-02	6.7502E-02	6.93108-02	7.1014E-02	7.5069E-02	8.2881E-02	9.2927E-02	0.1020	0.1073
5	9.4466E-02	9.5241E-02	9.4987E-02	9.1958E-02	8.6465E-02	8.1458E-02	8.1237E-02	8.6403E-02	9.2651E-02	9.6338E-02
4	0.1449	0.1366	0.1275	0.1181	0.1070	9.41508-02	8.3015E-02	7.94248-02	8.2387E-02	8.5477E-02
3	0.2283	0.1960	0.1659	0.1451	0.1288	0.1126	9.4428E-02	7.77432-02	7.0227E-02	/.0709E-02
Ż	0.4451	0.2723	0.2096	0.1702	0.1487	0.1326	0.1153	9.2307E-02	6.52038-02	4.7800E-02
ī	0.4916	0.1597	0.1205	9.2608E-02	8.1138E-02	7.5409E-02	7.0572E-02	6.2207E-02	4.5608E-02	1.7645E-02
					J = 2					
							•			
	1	2	3	4	5	6	7	8	9	10
11	2.12668-03	6.4845E-03	1.1234E-02	1.6811E-02	2.3920E-02	3.3821E-02	4.91458-02	7.6615E-02	0.1374	0.3114
10	9.3831E-03	1.53888-02	2.3889E-02	3.4464E-02	4.8098E-02	6.6772E-02	9.4014E-02	0.1364	0.2079	0.3609
	(					. /				
(						(				

9 1.7300E-02 2.1333E-02 2.8260E-02 3.7634E-02 4.9986E-02 6.6585E-02 8.9262E-02 0.1202 0.1611 0.2091 3.0016E-02 3.5810E-02 4.3767E-02 5.4269E-02 6.8151E-02 8.6094E-02 0.1079 8 2.6750E-02 0.1314 0.1508 7 4.0235E-02 4.3271E-02 4.7976E-02 5.3709E-02 6.1087E-02 7.1188E-02 8.4360E-02 9.9377E-02 0.1135 0.1230 7.1014E-02 7.5069E-02 8.2881E-02 9.2927E-02 0.1020 6 6.1574E-02 6.4471E-02 6.7502E-02 6.9310E-02 0.1073 5 9.4466E-02 9.5241E-02 9.4987E-02 9.1958E-02 8.6465E-02 8.1458E-02 8.1237E-02 8.6403E-02 9.2651E-02 9.6338E-02 0,1275 9.4150E-02 8.3015E-02 7.9424E-02 8.2387E-02 8.5477E-02 4 0.1449 0.1366 0.1181 0.1070 3 0.2283 0.1960 0.1659 0.1451 0.1288 0.1126 9.4428E-02 7.7743E-02 7.0227E-02 7.0709E-02 9.2307E-02 6.5203E-02 4.7800E-02 2 0.4451 0.2723 0.2096 0.1702 0.1487 0.1326 0.1153 0.1205 9.2608E-02 8.1138E-02 7.5409E-02 7.0572E-02 6.2207E-02 4.5608E-02 1.7645E-02 1 0.4916 0.1597

Y-DIRECTION - SOLUTE DISPERSIVE CONDUCTANCE BETWEEN Y(J) AND Y(J+1) (KG/S)

#### VERTICAL SLICES

J = 1

	1	2	3	4.	5	6	7	8	9	10
11	8.9629E-14	8.6104E-03	1.7718E-02	2.8044E-02	4.0729E-02	5.7738E-02	8.2963E-02	0.1258	0.2141	0.4488
10	8.3242E-03	2.3854E-02	3.8923E-02	5.8074E-02	8.2195E-02	0.1142	0.1593	0.2260	0.3274	0.4848
9	1.6646E-02	3.7577E-02	4.8806E-02	6.5119E-02	8.6613E-02	0.1150	0.1531	0.2042	0.2713	0.3596
8	2.6217E-02	5.5782E-02	6.4874E-02	7.8492E-02	9.6639E-02	0,1205	0.1518	0.1909	0.2366	0.2846
7	3.9872E-02	8.27632-02	9.0316E-02	0.1004	0.1132	0.1303	0.1537	0.1825	0.2131	0.2398
6	6.1793E-02	0.1258	0.1309	0.1353	0.1383	0.1437	0.1560	0.1753	0.1960	0.2118
-5	9.6675E-02	0.1900	0.1881	0.1847	0.1762	0.1652	0.1599	0.1666	0.1802	0.1911
4	0.1531	0.2812	0.2589	0.2417	0.2229	0.1991	0.1741	0.1597	0.1620	0.1699
3	0.2499	0.4194	0,3483	0.3052	0.2719	0.2404	0.2056	0.1693	0.1454	0.1418
2	0.6396	0.5467	0.4702	0.3757	0.3181	0.2814	0.2481	0.2072	0.1551	0.1085
1	0.0000	0.6513	0.2802	0.2132	0.1739	0.1568	0.1463	0.1331	0.1079	6.2509E-02

11 11 0.0000 10 0.4048 9 0.2215 8 0.1541 7 0.1238 6 0.1072 5 9.5650E-02 4 8.4157E-02 3 6.8402E-02 2 4.3339E-02 1 0.0000

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# Z-DIRECTION - SOLUTE DISPERSIVE CONDUCTANCE BETWEEN Z(K) AND Z(K+1) (KG/S)

#### VERTICAL SLICES

#### J = 1

	1	2	3	4	5	6	7	8	9	10
10	2.1119E-03	9.5084E-03	1.8033E-02	2.8037E-02	4.0362E-02	5.6722E-02	8.0326E-02	0.1180	0.1880	0.3531
9	6.2519E-03	1.51876-02	2.1671E-02	3.0550E-02	4.1994E-02	5.7195E-02	7.8294E-02	0.1087	0.1532	0.2158
8	1.0557E-02	2.3064E-02	2.8270E-02	3.5924E-02		5.9359E-02			0.1284	0.1590
7	1.5982E-02	3.3803E-02	3.8482E-02	4.4930E-02	5.3074E-02	6.3667E-02	7.7490E-02	9.4400E-02	0.1127	0.1290
6	2.4132E-02	5.0224E-02	5.4470E-02	5.8951E-02	6.35538-02	6.9706E-02	7.8804E-02	9.0508E-02	0.1025	0.1118
5	3.7475E-02	7.6146E-02	7.8616E-02	7.9678E-02	7,8757E-02	7.7858E-02	8.01908-02	8.6772E-02	9.47088-02	0.1005
4	5.8807E-02	0.1146	0.1112	0.1066	9.9591E-02	9.0746E-02	8.3505E-02	8.2480E-02	8.6751E-02	9.0914E-02
3	9.4083E-02	0.1711	0.1532	0.1377	0.1237	0.1094	9.39658-02	8.1496E-02	7.76888-02	7.9561E-02
2	0.1557	0.2634	0.2095	0.1713	0.1475	0.1301	0.1127	9.26138-02	7.36208-02	6.4279E-02
1	0.4838	0.4533	0.2580	0.2000	0.1661	0.1485	0.1349	0.1177	9.08438-02	5.4444E-02

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11 10 0.3147 9 0.1467 8 9.1353E-02 7 6.8849E-02 6 5.7832E-02 5 5.1310E-02 4 4.6200E-02 3 4.0449E-02 2 3.1819E-02 1 1.6073E-02

J = 2

	- 1	2	3	4	•	-	•	8	9	10
10	2.1119E-03	9.5084E-03	1.8033E-02	2.8037E-02	4.0362E-02	5.6722E-02	8.03268-02	0.1180	0.1880	0.3531
9	6.2519E-03	1.5187E-02	2.1671E-02	3.0550E-02	4.1994E-02	5.71958-02	7.8294E-02	0.1087	0 1532	0.2158
8	1.0557E-02	2.3064E-02	2.8270E-02	3.5924E-02	4.6032E-02	5.93598-02	7.7066E-02	0.1001	0.1284	0.1590
7	1.5982E-02	3.3803E-02	3.8482E-02	4.4930E-02	5.3074E-02	6.3667E-02	7.7490E-02	9.4400E-02	0.1127	0.1290
6	2.41328-02	5.0224E-02	5.4470E-02	5.8951E-02	6.3553E-02	6.9706L-02	7.8804E-02	9.0508E-02	0,1025	0.1118
5	3.7475E-02	7.6146E-02	7.8616E-02	7.9678E-02	7.8757E-02	7.7858E-02	8.01908-02	8.6772L-02	9.4/081-02	0.1005
4	5.8807E-02	0.1146	0.1112	0.1066	9.9591E-02	9.07468-02	8.3505E-02	8.2480E-02	8.6/51L-02	9.0914E-02

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	2	9.4083E-02 0.1557 0.4838	0.1711 0.2634 0.4533	0.1532 0.2095 0.2580	0.1377 0.1713 0.2000	0.1237 0.1475 0.1661	0.1094 0.1301 0.1485	9.3965E-02 0.1127 0.1349		7.7688E-02 7.3620E-02 9.0843E-02	6.4279E-02	
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	11	
10	0.3147	
9	0.1467	
8	9.1353E-02	
7	6.8849E-02	
6	5.7832E-02	
5	5.1310E-02	 
- 4	4.6200E-02	• : .
3	4.0449E-02	
2	3.1819E-02	
1	1.6073E-02	

X-DIRECTION - INTERSTITIAL PORE VELOCITY BETWEEN X(I) AND X(I+1) ( M/S)

VERTICAL SLICES

	1	2	3	4	5	6	7	8	9	10
11	5.31478-03	1.6203E-02	2.8062E-02	4,1978E-02	5.9706E-02	8.4383E-02	0.1226	0.1910	0.3425	0.7758
10	5.11938-03	1.5614E-02	2.6965E-02	4.0079E-02	5.6353E-02	7.7922E-02	0.1080	0.1502	0.2024	0.2096
ĝ	4.8136E-03	1.4572E-02	2.4790E-02	3.6012E-02	4.9015E-02	6.4614E-02	8,2953E-02	0.1013	0.1079	6.9241E-02
8	4.9878E-03	1.4765E-02	2.4096E-02	3.3120E-02	4.2236E-02	5.1673E-02	6.0621E-02	6.5697E-02	5.8997E-02	2.8473E-02
7	6.90048-03	1.9542E-02	2.94708-02	3.6502E-02	4.1330E-02	4.4676E-02	4.6368E-02	4.4429E-02	3.4949E-02	1.434?F-02
6	1.19386-02	3.2313E-02	4.5418E-J2	5.1154E-02	5.1235E-02	4.7783E-02	4.2351E-02	3.4°C0E-02	<b>2.</b> 4216E-02	8.9491E-03
5	2.0321E-02	5.24598-02	7.05998-02	7.62888-02	7.2688E-02	6.2804E-02	4.96498-02	3.5753E-02	2.1836E-02	7.4139E-03
4	3.5123E-02	8.1111E-02	0.1016	0.1056	9.9527E-02	8.6266E-02	6.7538E-02	4.6358E-02	2.6284E-02	
3	7.2376E-02	0.1316	0.1430	0.1368	0.1258	0.1114	9.2011E-02	6.6718E-02	3.8457E-02	1.2019E-02
2	0.1688	0.2351	0.1988	0.1674	0.1485	0.1340	0.1183	9.5473E-02	6.1827E-02	2.0481E-02
1	0.9832	0.3194	0.2411	0.1854	0.1630	0.1527	0.1452	0.1316	0.1008	4.1060E-02
				1. A.	) _ <b>1</b>					
	J = 2									
	1	2	3	4	5	6	7	8	9	10
11	5.3147E-03	1.6203E-02	2.8062E-02	4.1978E-02	5.9706E-02	8.4383E-02	0.1226	0.1910	0.3425	0.7758
10	5.1193E-03	1.5614E-02	2.6965E-02	4.0079E-02	5.6353E-02	7.7922E-02	0.1080	0.1502	0.2024	0.2096
9	4.8136E-03	1.4572E-02	2.4790E-02	3.6012E-02	4.9015E-02	6.4614E-02	8.2953E-02	0.1013	0.1079	6.9241E-02

8	4.9878E-03	1.4765E-02	2.4096E-02	3.3120E-02	4.22362-02	5.1673E-02	6.0621E-02	6.5697E-02	5.8997E-02	2.8473E-02
7	6.9004E-03		2.9470E-02							
6	1.1938E-02	3.2313E-02	4,5418E-02	5.1154E-02	5.1235E-02	4.7783E-02	4.2351E-02	3.4980E-02	2.4216E-02	8.9491E-03
- 5	2.0321E-02	5.2459E-02	7.0599E-02	7.6288E-02	7.26888-02	6.2804E-02	4.9649E-02	3.5753E-02	2.1836E-02	7.4139E-03
- 4	3.5123E-02	8.1111E-02	0.1016	0.1056	9.9527E-02	8.6266E-02	6.7538E-02	4.6358E-02	2.62842-02	8.3966E-03
3	7.2376E-02	0.1316 👘	0.1430	0.1368	0.1258	0.1114	9.2011E-02	6.6718E-02	3.8457E-02	1.20198-02
2	0.1688	0.2351	0.1988	0.1674	0.1485	0.1340	0.1183	9.5473E-02	6.1827E-02	2.0481E-02
1	0.9832	0.3194	0.2411	0.1854	0.1630	0.1527	0.1452	0.1316	0.1008	4.1060E-02

Y-DIRECTION - INTERSTITIAL PORE VELOCITY BETWEEN Y(J) AND Y(J+1) ( H/S)

VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
- 11	-2.2402E-13	1.3446E-13	-2.0182E-13	2.0198E-13	-1.7970E-13	5.62168-14	-7.8794E-14	5.6347E-14	-4.5119E-14	1.6927E-14
10	1.8026E-13	-1.3525E-13	2.7075E-13	-2.0330E-13	9.0463E-14	-1.5849E-13	4.5321E-14	-9.0667E-14	4.5300E-14	0.0000
	-1.3981E-13									
8	2.5748E-13				1.4991E-13				9.3202E-14	-9.1674E-14
7	0.0000		-2.3982E-13						0.0000	4.1696E-13
			2.8588E-13							-9.4088E-14
5	-4.0666E-13	3.2358E-13	-5.5724E-13	7.7364E-14	-7.40208-14	1.3884E-13	-2.5461E-13	5.7417E-14	-3.1115E-13	1.9227E-13
- 4	3.3193E-13	-6.6295E-13	4.9458E-13	-1.6275E-13	3.1729E-13	-4.5482E-13	1.4059E-13	-6.2883E-14	1.1038E-13	-2.9697E-13
3	0.0000	8.3124E-13	-1.6604E-13	4.9585E-13	-1.6319E-13					
2	1.6631E-13	-1.6630E-13	0.0000	0.0000	4.9451E-13	-8.0921E-13	-3.1128E-13	-2.8970E-13	1.2848E-13	-1.0865E-13
1	0.0000	4.9892E-13	0.000	1.6609E-13	-4.9572E-13	0.0000	-4.7197E-13	0.0000	-1.3341E-13	3.4561E-13

- 11 11 0.0000 10 3.3909E-14 9 2.2678E-14 8 0.0000 7 -2.7487E-13 6 1.8491E-13 5 -4.6734E-14 4 3.3182E-13 3 0.0000 2 2.9379E-13 1 0.0000

# Z-DIRECTION - INTERSTITIAL PORE VELOCITY BETWEEN Z(K) AND Z(K+1) ( M/S)

VERTICAL SLICES

] = 1	
-------	--

	1	2	3	1. The <b>4</b> - 1. Th	5	6	7	8	9 ·	10
10	5.2721E-03	5.4081E-03	5.9016E-03	6.9400E-03	8.8591E-03	· 1.2351E-02	1.91238-02	3.4269E-02	7.5822E-02	0.2168
9	1.5484E-02	1.5907E-02	1.7325E-02	2.0207E-02	2,5363E-02	3.4215E-02	4.9494E-02	7.6856E-02	0.1282	0.2243
8	2.5468E-02	2.6055E-02	2.8004E-02	3.1947E-02	3.8890E-02	5.0277E-02	6.8156E-02	9.53128-02	0.1348	0.1858
7	3.6571E-02	3.6981E-02	3.8489E-02	4.2033E-02	4.8865E-02	6.0284E-02	7.7367E-02	0.1004	0.1282	0.1557
6	5.1752E-02	5.1034E-02					7.9208E-02			0.1358
-		7.2393E-02					7.3757E-02			
4	0.1181	0.1050	8.2061E-02	6.2301E-02	5.1263E-02	5.0606E-02	6.0307E-02	·7.7522E-02	9.5784E-02	0.1087
. 3	0.1892	0.1513					4.0916E-02			
-	0.3115	0.2152					2.0668E-02			
1	0.9677	0.1533	6.9135E-02	2.7564E-02	1.2346E-02	5.8178E-03	3.9359E-03	6.3626E-03	1.4191E-02	2.7546E-02

$\begin{array}{cccccccccccccccccccccccccccccccccccc$	10 9 8 7 6 5 4 2	11 0.7839 0.3651 0.2270 0.1707 0.1430 0.1264 0.1132 9.8536E-02					• •				
J = 2 $J = 2$ $1  2  3  4  5  6  7  8  9  10$	3										
J = 2 1 2 3 4 5 6 7 8 9 10	1										
1 2 3 4 5 6 7 8 9 10	-	3103/32-02									
1 2 3 4 5 6 7 8 9 10						J = 2	•				
en la la culta de la culta d											
10 5.2721E-03 5.4081E-03 5.9016E-03 6.9400E-03 8.8591E-03 1.2351E-02 1.9123E-02 3.4269E-02 7.5822E-02 0.2168		1	2	3	4	5	6	7	8	9	10
	10	5.2721E-03	5.4081E-03	5.9016E-03	6.9400E-03	8.8591E-03	1.2351E-02	1.9123E-02	3.4269E-02	7.5822E-02	0.2168
9 1.5484E-02 1.5907E-02 1.7325E-02 2.0207E-02 2.5363E-02 3.4215E-02 4.9494E-02 7.6856E-02 0.1282 0.2243	9	1.5484E-02	1.5907E-02	1.7325E-02	2.0207E-02	2.5363E-02	3.4215E-02	4.9494E-02	7.6856E-02	0.1282	0.2243
8 2.5468E-02 2.6055E-02 2.8004E-02 3.1947E-02 3.8890E-02 5.0277E-02 6.8156E-02 9.5312E-02 0.1348 0.1858	8	2.5468E-02	2.6055E-02	2.8004E-02	3.1947E-02	3.8890E-02	5.0277E-02	6.8156E-02	9.5312E-02	0.1348	0.1858
7 3.6571E-02 3.6981E-02 3.8489E-02 4.2033E-02 4.8865E-02 6.0284E-02 7.7367E-02 0.1004 0.1282 0.1557	7	3.6571E-02	3.6981E-02	3.8489E-02	4.2033E-02	4.8865E-02	6.0284E-02	7.7367E-02	0.1004	0.1282	0.1557
6 5.1752E-02 5.1034E-02 4.9809E-02 5.0282E-02 5.4568E-02 6.4093E-02 7.9208E-02 9.8604E-02 0.1191 0.1358	6	5.1752E-02	5.1034E-02	4.9809E-02	5.0282E-02	5.4568E-02	6.4093E-02	7.9208E-02	9.8604E-02	0.1191	0.1358
5 7.6668E-02 7.2393E-02 6.3791E-02 5.6736E-02 5.5122E-02 6.0807E-02 7.3757E-02 9.1384E-02 0.1089 0.1217	5	7.66688-02	7.2393E-02	6.3791E-02	5.6736E-02	5.5122E-02	6.0807E-02	7.3757E-02	9.1384E-02	0.1089	0.1217

7	3.6571E-02	3.6981E-02	3.8489E-02	4.2033E-02	4.8865E-02	6.0284E-02	7.7367E-02	0.1004	0.1282	0.1557
6	5.1752E-02	5.1034E-02	4.9809E-02	5.0282E-02	5.4568E-02	6.4093E-02	7.9208E-02	9.8604E-02	0.1191	0.1358
5	7.66688-02	7.2393E-02	6.3791E-02	5.6736E-02	5.5122E-02	6.0807E-02	7.3757E-02	9.1384E-02	0.1089	0.1217
4	0.1181	0.1050	8.2061E-02	6.2301E-02	5.1263E-02	5.0606E-02	6.0307E-02	7.7522E-02	9.5784E-02	0.1087
3	0.1882	0.1513	0.1024	6.5566E-02	4.4378E-02	3.6571E-02	4.0916E-02	5.6056E-02	7.6291E-02	9.2439E-02

2	0.3115	0.2152	0.1120	5.7847E-02	3.2313E-02	2.1230E-02	2.0668E-02	3.0104E-02	4.8174E-02	6.7575E-02	
1	0.9677	0.1533	6.9135E-02	2.7564E-02	1.2346E-02	5.8178E-03	3.9359E-03	6.3626E-03	1.4191E-02	2.7546E-02	
7	11 0.7839 0.3651 0.2270 0.1707 0.1430 0.1264 0.1132 9.8536E-02 7.7023E-02 3.8579E-02										

*** OUTPUT AT END OF TIME STEP NO. 50 ***			
TINE	10.0	(S);	1.157E-04 (D)
CURRENT TIME STEP LENGTH	0.200	(5);	2.315E-06 (D)
NO. OF P.T.C LOOP ITERATIONS USED	2 4.6795E+ 2.1016E-	02 ( PA) 02 (-)	AT LOCATION ( 2. 2. 1) AT LOCATION (10, 2, 1)

PRESSURE ( PA)

VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
11	4586.2	4571.1	4525.2	4445.8	4327.7	4160.5	3925.9	3587.1	3062.0	2123.5
10	6172.7	6158.3	6114.2	6038.3	5925.9	5768.9	5553.2	5256.2	4844.6	4291.3
9	7800.5	7787.3	7747.1	7678.9	7580.0	7446.2	7270.5	7045.8	6771.4	6478.4
8	9494.4	9481.8	9444.3	9382.7	9297.4	9187.9	9053.1	8893.9	8719.4	8560.8
7	11292.	11278.	11236.	11171.	11089.	10992.	10885.	10769.	10655.	10563.
6	13217.	13196.	13139.	13058.	12961.	12859.	12757.	12661.	12576.	12514.
5	15253.	15221.	15139.	15028.	14905.	14781.	14665.	14565.	14485.	14432.
4	17378.	17323.	17201.	17050.	16893.	16740.	16599.	16478.	16385.	16325.
	1									

Spec. at

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 $\sim 10^{10}$ .....

321	19619. 22135. 25000.	19496. 21737. 24092.	19298. 21407. 23468.	19093. 21131. 23130.	18896. 20894. 22867.	18713. 20682. 22637.	18544. 20487. 22417.	18395. 20305. 22199.	16273. 20144. 21985.	18192. 20023. 21797.
11 10 9 8 7 6	11 0.00000 3718.6 6289.9 8483.6 10524. 12490.	•								
· 5	14413. 16305.	• •						•		
3	18164.									
2	19977.									
1	21709.	· . ·						2 2		
	•				<b>)</b> =	2				
11 10 9 8 7 6 5 4 3 2 1	1 4586.2 6172.7 7800.5 9494.4 11292. 13217. 15253. 17378. 19619. 22135. 25000.	2 4571.1 6158.3 7787.3 9481.8 11278. 13196. 15221. 17323. 19496. 21737. 24092.	3 4525.2 6114.2 7747.1 9444.3 11236. 13139. 15139. 17201. 19298. 21407. 23468.	4 4445.8 6038.3 7678.9 9382.7 11171. 13058. 15028. 17050. 19093. 21131. 23130.	5 4327.7 5925.9 7580.0 9297.4 11089. 12961. 14905. 16893. 18896. 20894. 22867.	6 4160.5 5768.9 7446.2 9187.9 10992. 12859. 14781. 16740. 18713. 20682. 22637.	7 3925.9 5553.2 7270.5 9053.1 10885. 12757. 14665. 16599. 18544. 20487. 22417.	8 3587.1 5256.2 7045.8 8893.9 10769. 12661. 14565. 16478. 18395. 20305. 22199.	9 3062.0 4844.6 6771.4 8719.4 10655. 12576. 14485. 16385. 18273. 20144. 21985.	10 2123.5 4291.3 6478.4 8560.8 10563. 12514. 14432. 16325. 18192. 20023. 21797.

11 11 0.00000 10 3718.6 9 6289.9 8 8483.6 7 10524.

6 12490. 5 14413. 4 16305.

3 18164. 2 19977. 1 21709.

288

SOLUTE SCALED MASS FRACTION (-)

VERTICAL SLICES

J = 1. 1 0.00111 2 0.00179 5 3 0.00293 7 10 4 6 8 9 0.00435 0.00603 0.01016 0.01238 0.01420 0.01515 11 0.00798 0.01133 0.01367 0.01199 0.01577 0.02012 0.02186 0.02258 10 0.01798 0.02151 0.01895 0.06952 0.22294 0.48762 0.06912 0.23719 0.06946 0.06639 0.06180 0.05359 0.02927 9 0.06924 0.06861 0.04161 0.20745 0.44239 0.70692 0.89107 8 0.23339 0.18837 0.16590 0.13891 0.10664 0.07184 0.04297 0.25755 0.41289 0.58674 0.74331 0.84745 0.89977 0.91610 0.53015 0.51866 0.38761 0.32582 0.52739 7 0.18448 0.11436 0.06186 0.76825 0.93480 0.98714 0.99770 0.99951 0.82292 0.80818 0.62622 6 0.28967 0.17356 0.08869 0.95923 0.82304 0.72342 0.12594 5 0.96737 0.42091 0.25350 0.17596 0.24235 0.33220 0.42791 43 0.96779 0.99088 0.92904 0.85975 0.92938 0.99701 0.99501 0.56766 0.35659 0.70243 0.78922 0.99978 0.99948 0.97181 0.48055 0.99994 0.99693 0.95926 0.60195 2 0.99999 0.98686 0.99978 0.99050 1 1.00000 0.99999 0.99813 0.96757 0.81989 0.66059

11 10 9 8 7 6 5 4 3 2 1	11 0.01601 0.01823 0.02413 0.03158 0.04223 0.05746 0.05746 0.10288 0.10288 0.13074 0.15963 0.20642				J = 2					
11 10	1 0.00111 0.01133	2 0.00179 0.01199	3 0.00293 0.01367	4 0.00435 0.01577	5 0.00603 0.01798	6 0.00798 0.02012	7 0.01016 0.02186	8 0.01238 0.02258	9 0.01420 0.02151	10 0.01515 0.01895

9 8 7 6 5 4 3 2 1	0.06912 0.23719 0.53015 0.82292 0.96737 0.99701 0.99978 0.99999 1.00000	0.06924 0.23339 0.51866 0.80818 0.95923 0.99501 0.99948 0.99994 0.99999	0.06952 0.22294 0.48762 0.76825 0.93480 0.98714 0.99770 0.99951 0.99978	0.06946 0.20745 0.44239 0.70692 0.89107 0.96779 0.99088 0.99683 0.99813	0.06861 0.18837 0.38761 0.62622 0.82304 0.92904 0.97181 0.98686 0.99050	0.06639 0.16590 0.32582 0.52739 0.72342 0.85975 0.92938 0.95926 0.96757	0.06180 0.13891 0.25755 0.41289 0.58674 0.74331 0.84746 0.89977 0.91610	0.05359 0.10664 0.18448 0.28967 0.42091 0.56766 0.70243 0.78922 0.81989	0.04161 0.07184 0.11436 0.25350 0.35659 0.48055 0.60195 0.66059	0.02927 0.04297 0.06186 0.08869 0.12594 0.17596 0.24235 0.33220 0.42791
1	1.00000	0.999999	0,99978	0.99813	0.99050	0.96757	0.91610	0.81989	0.66059	0.42791

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• •	11
11	0.01601
10	0.01823
. 9	0.02413
.8	0.03158
7	0.04223
б	0.05746
5	0.07785
4 .	0.10288,
3	0.13074
2	0.15963
1	0.20642

# DENSITY (KG/ M\*\*3)

VERTICAL SLICES

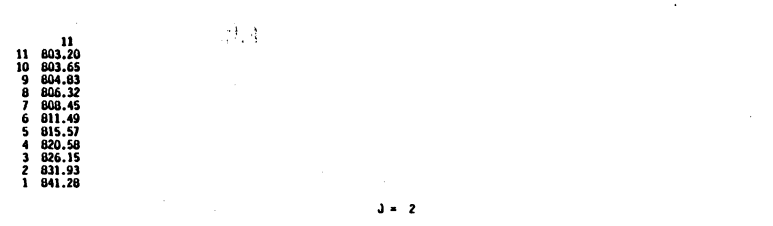
J = 1

	1	2	3	4	5	6	7	8	9	10
11	800.22	800.36	800.59	800.87	801.21	801.60	802.03	802.48	802.84	803.03
10	802.27	802.40	802.73	803.15	803.60	804.02	804.37	804.52	804.30	803.79
9	813.82	813.85	813.90	813.89	813.72	813.28	812.36	810.72	808.32	805.85
8	847.44	846.68	844.59	841.49	837.67	833.18	827.78	821.33	814.37	808.59
7	906.03	903.73	897.52	888.48	877.52	865.16	851.51	836.90	822.87	812.37
6	964.58	961.64	953.65	941.38	925.24	905.48	882.58	857.93	834.71	817.74
5	993.47	991.85	986.96	978.21	964.61	944.68	917.35	884.18	850.70	825.19
4	999.40	999.00	997.43	993.56	985.81	971.95	948.66	913.53	871.32	835.19
3	999.96	999.90	999.54	998.18	994.36	985.88	969.49	940.49	896.11	848.47
2	1000.0	999.99	999.90	999.39	997.37	991.85	979.95	957.84	920.39	866.44
1	1000.0	1000.0	999.96	999.63	998.10	993.51	983.22	963.98	932.12	885.58

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	1	2	3	4	5	6	7	8	9	10
11	800.22	800.36	800.59	800.87	801.21	801.60	802.03	802.48	802.84	803.03
10	802.27	802.40	802.73	803.15	803.60	804.02	804.37	804.52	804.30	803.79
9	813.82	813.85	813.90	813.89	813.72	813.28	812.36	810.72	808.32	805.85
8	847.44	846.68	844.59	841.49	837.67	833.18	827.78	821.33	814.37	808.59
7	906.03	903.73	897.52	888.48	877.52	865.16	851.51	836.90	822.87	812.37
6	964.58	961.64	953.65	941.38	925.24	905.48	882.58	857.93	834.71	817.74
5	993.47	991.85	986.96	978.21	964.61	944.68	917.35	884.18	850.70	825.19
4	999.40	999.00	997.43	993.56	985.81	971.95	948.66	913.53	871.32	835.19
3	999.96	999.90	999.54	998.18	994.36	985.88	969.49	940.49	896.11	848.47
2	1000.0	999.99	999.90	999.39	997.37	991.85	979.95	957.84	920.39	866.44
1	1000.0	1000.0	999.96	999.63	998.10	993.51	983.22	963.98	932.12	885.58

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- 11 11 803.20 10 803.65 9 804.83 8 806.32 7 808.45 6 811.49 5 815.57 4 820.58 3 826.15 2 831.93 1 841.28

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#### VISCOSITY (KG/ M-S)

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VERTICAL SLICES

										• .
					J = 1					
		and the second second								
	1	2	3	1 <b>4</b> 1	5	6	7	8	9	10
11	1.2991E-03	1.2986E-03	1.2976E-03	1.2965E-03	1.2952E-03	1.2936E-03	1.29198-03	1.2901E-03	1.2886E-03	1.2879E-03
10	1.2909E-03	1.2904E-03	1.2890E-03	1.2874E-03	1.2856E-03	1.2839E-03	1.2825E-03	1.2820E-03	1.2828E-03	1.2848E-03
9	1.2455E-03	1.2455E-03	1.2452E-03	1.2453E-03	1.2459E-03	1.2477E-03	1.2512E-03	1.2576E-03	1.2669E-03	1.2767E-03
8	1.1225E-03	1.1251E-03	1.1324E-03	1.1433E-03	1.1569E-03	1.1731E-03	1.1929E-03	1.2170E-03	1.2435E-03	1.2659E-03
7	9.3630E-04	9.42998-04	9.6128E-04	9.88572-04	1.0227E-03	1.0625E-03	1.1084E-03	1.1597E-03	1.2111E-03	1.2512E-03
6	7.8110E-04	7.88262-04	8.0799E-04	8.3925E-04	8.8224E-04	9.3790E-04	1.0068E-03	1.0866E-03	1.1676E-03	1.2305E-03
· 5	7.1428E-04	7.1789E-04	7.28832-04	7.4883E-04	7.8104E-04	8.3072E-04	9.0407E-04	1.0018E-03	1.1112E-03	1.2025E-03
4	7.0129E-04	7.0216E-04	7.0560E-04	7.1410E-04	7.3144E-04	7.63492-04	8.2055E-04	9.1481E-04	1.0425E-03	1.1658E-03
3	7.0009E-04	7.00238-04	7.0100E-04	7.0396E-04	7.1232E-04	7.3128E-04	7.69328-04	8.41598-04	9.6550E-04	1.1189E-03
<u>ž</u>	7.0001E-04	7.0003E-04	7.00218-04	7.0133E-04	7.0572E-04	7.1788E-04	7.4481E-04	7.9756E-04	8.95598-04	1.0584E-03
Ī	7.0000E-04	7.0001E-04	7.0010E-04	7.0081E-04	7.0413E-04	7.1420E-04	7.3732E-04	7.8256E-04	8.6367E-04	9.9747E-04
	11									
~ 11	1.2872E-03									
10										

10	1.20346-03				
9	1.2807E-03				
8	1.2748E-03				•
7	1.2665E-03				
6	1.2546E-03				
5		A LONG AND A REAL PROPERTY OF			· · · · ·
4	1.2198E-03	•	e • ja se en la se en	•	
3	1.1989E-03	· · · · · · · · · · · · · · · · · · ·	•		
2	1.1777E-03				
ī	1.1441E-03				
•	9 9 9 1 1 1 4 W MA	•			

J = 2

	1	2	3	- 4	5	6	7	8	9	10
11	1.2991E-03	1.2986E-03	1.2976E-03	1.2965E-03	1.2952E-03	1.2936E-03	1.2919E-03	1.2901E-03	1,2886E-03	1.2879E-03
10	1.2909E-03	1.2904E-03	1.2890E-03	1.2874E-03	1.2856E-03	1.2839E-03	1.2825E-03	1.2820E-03	1.2828E-03	1.2848E-03
9	1.2455E-03	1.2455E-03	1,2452E-03	1.2453E-03	1.2459E-03	1.2477E-03	1.2512E-03	1.2576E-03	1.2669E-03	1.2767E-03
8	1.1225E-03	1.1251E-03	1.1324E-03	1.1433E-03	1.1569E-03	1.1731E-03	1.1929E-03	1.2170E-03	1.2435E-03	1.2659E-03
7	9.3630E-04	9.4299E-04	9.6128E-04	9.8857E-04	1.0227E-03	1.0625E-03	1.1084E-03	1.1597E-03	1.2111E-03	1.2512E-03
							1			

6	7.8110E-04	7.8826E-04	8.0799E-04	8.3925E-04	8.8224E-04	9.3790E-04	1.0068E-03	1.0866E-03	1.1676E-03	1.2305E-03
5	7.1428E-04	7.1789E-04	7.2883E-04	7.4883E-04	7.8104E-04	8.3072E-04	9.0407E-04	1.0018E-03	1.1112E-03	1.2025E-03
4	7.01298-04	7.0216E-04	7.0560E-04	7.14102-04	7.3144E-04	7.6349E-04	8.20558-04	9.1481E-04	1.0425E-03	1.1658E-03
3	7.0009E-04	7.0023E-04	7.0100E-04	7.0396E-04	7.1232E-04	7.3128E-04	7.6932E-04	8.4159E-04	9.6550E-04	1.1189E-03
2	7.00018-04	7.0003E-04	7.0071E-04	7.0133E-04	7.0572E-04	7.1788E-04	7.4481E-04	7.9756E-04	8.9559E-04	1.0584E-03
1	7.0000E-04	7.0001E-04	7.0010E-04	7.0081E-04	7.0413E-04	7.1420E-04	7.3732E-04	7.8256E-04	8.6367E-04	9.9747E-04

- 11 11 1.2872E-03 10 1.2854E-03 9 1.2807E-03 8 1.2748E-03 7 1.2665E-03 6 1.2546E-03 5 1.2388E-03 5 1.2198E-03 3 1.1989E-03 2 1.1777E-03 1 1.1441E-03

292

*** GLOBAL FLOW BALANCE SUNNARY *** CURRENT TIME STEP	RATES	·	AMOUNTS	
FLUID INFLOW FLUID OUTFLOW CHANGE IN FLUID IN REGION RESIDUAL IMBALANCE FRACTIONAL IMBALANCE	2.580780E+00 6.645022E-01 -5.047900E-06	(KG/S) (KG/S) (KG/S) (KG/S)	6.490574E-01 5.161559E-01 1.329004E-01 -1.009580E-06 0.0000	(KG) (KG) (KG) (KG)
SOLUTE INFLOW SOLUTE OUTFLOW CHANGE IN SOLUTE IN REGION	1.915943E-04	(KG/S) (KG/S) (KG/S)	3.245291E-03 3.831887E-05 3.206945E-03	(KG) (KG) (KG)
RESIDUAL IMBALANCE		(KG/S)	-2.733812E-08 0.0000	(KG)

#### CUHULATIVE SUNNARY

AMOUNTS

FLUID INFLOW	3.325127E+01	(KG)
FLUID OUTFLOW	2.626598E+01	(KG)

......

CHANGE IN FLUID IN REGION		(KG) (KG)
RESIDUAL INBALANCE	-2.603850E-04	(KG)
SOLUTE INFLOW	1.665237E-01	(KG)
SOLUTE OUTFLOW		(KG)
CHANGE IN SOLUTE IN REGION	1.663889E-01	(KG)
SOLUTE IN REGION		(KG)
RESIDUAL INBALANCE	9.057561E-05	(KG)
FRACTIONAL IMBALANCE		· -/
CIMMU ATTUE COECIETED & CELL CLUID NET INCLUI	6 0052065.00	(20)

COMOLATIVE SPECIFIED P CELL FLUID NET INFLOW	0.9832806+00	(Ku)
CUMULATIVE FLUX B.C. FLUID NET INFLOW	0.000000E-01	(KG)
CUMULATIVE LEAKAGE B.C. FLUID NET INFLOW	0.000000E-01	ĊKGŚ
CUMULATIVE AQUIFER INFLUENCE FLUID NET INFLOW	0.00000E-01	(KG)

#### CUMULATIVE SPECIFIED C CELL OR ASSOCIATED WITH

SPECIFIED P CELL SOLUTE HET INFLOW	1.662984E-01	(KG)
CUMULATIVE FLUX B.C. SOLUTE NET INFLOW	0.000000E-01	(KG)
CUMULATIVE LEAKAGE B.C. SOLUTE NET INFLOW		
CUMULATIVE AQUIFER INFLUENCE SOLUTE NET INFLOW		

#### SPECIFIED PRESSURE, TEMPERATURE, OR MASS FRACTION B.C. FLOW RATES POSITIVE IS INTO THE REGION

FLUID (KG/S)

VERTICAL SLICES

#### J = 1

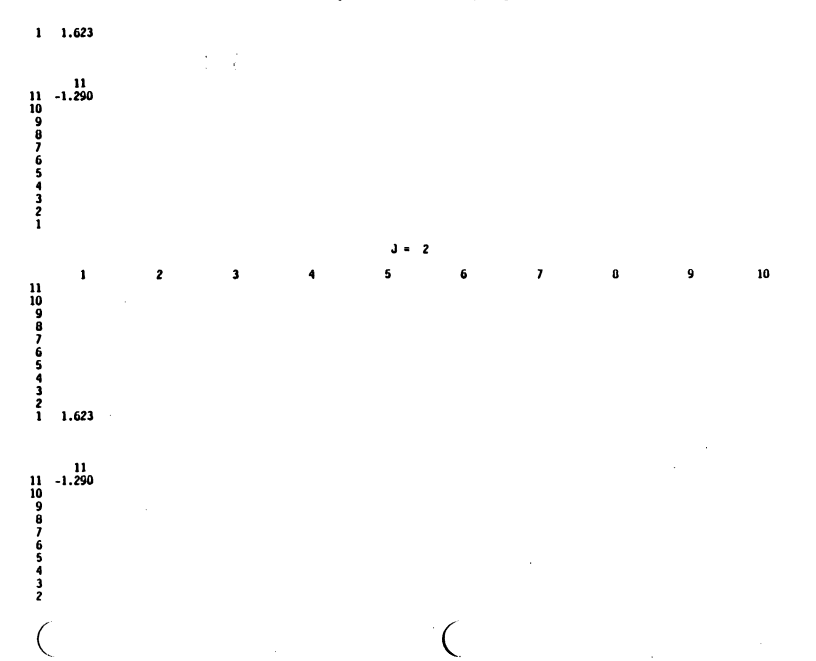
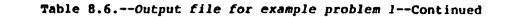
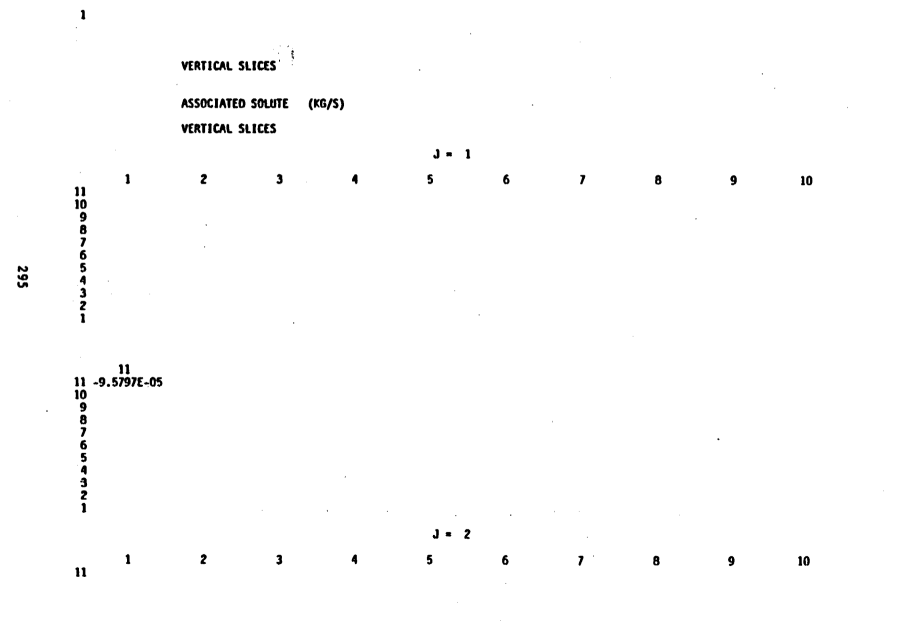


Table 8.6.--Output file for example problem 1--Continued



1.00



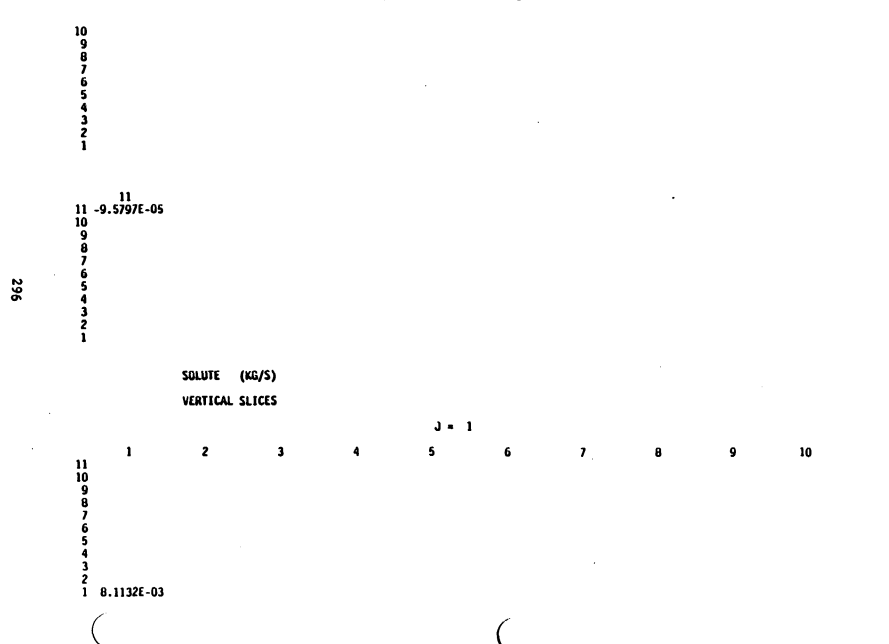
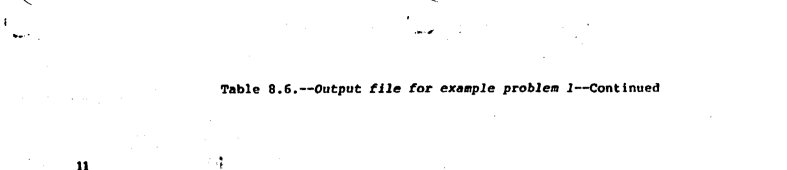


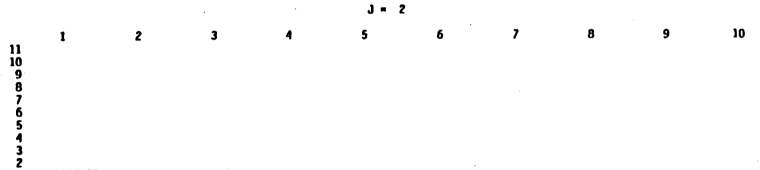
Table 8.6.--Output file for example problem 1--Continued



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10 9 8 7

5 



1 8.1132E-03

10 2 1

)

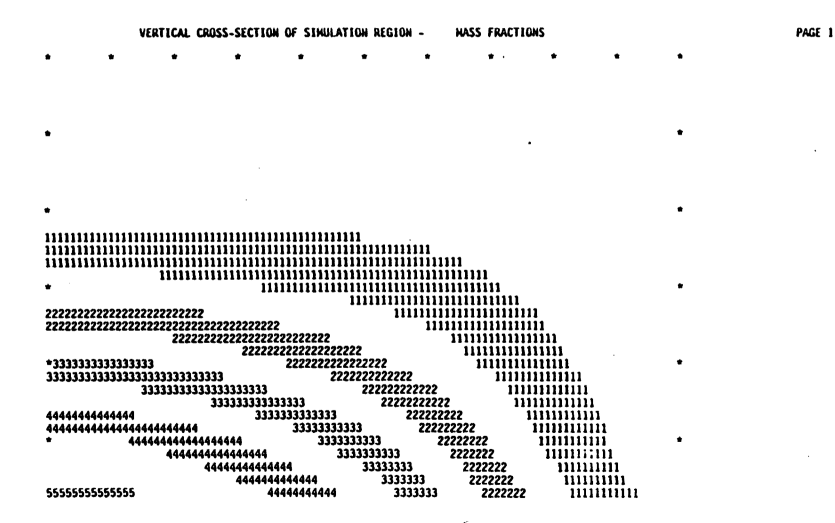


Table 8.6.--Output file for example problem 1--Continued

VERTICAL PLANE AT ROW NO. 1

. .

555555555555555555555555555555555555555	444444444	3333333	2222222	
*55555555555555555555555555555555555555	44444444	333333	222222	11111111111 +
555555555555555555555555555555555555555	44444444	333333	222222	1111111111
555555555555555555555555555555555555555	44444444	333333	222222	11111111111
555555555555555555555555555555555555555	444444	44 333333	222222	11111111111
555555555555555555555555555555555555555	55 4444	4444 3333	3 22222	111111111111
555555555555555555555555555555555555555	55555 44	44444 333	333 22222	1111111111
*55555555555555555555555555555555555555			3333 22222	2 11111111111
555555555555555555555555555555555555555	555555555	444444	33333 2222	22 1111111111
555555555555555555555555555555555555555	55555555555	444444	33333 222	22 1111111111
555555555555555555555555555555555555555	5555555555555	44444		222 11111111
555555555555555555555555555555555555555	555555555555555	444444		2222 11111111
555555555555555555555555555555555555555	555555555555555555555555555555555555555	444444		1111111
*55555555555555555555555555555555555555		444444		2222 111111*
555555555555555555555555555555555555555	555555555555555555555555555555555555555	44444		2222 111111
555555555555555555555555555555555555555				2222 11111
555555555555555555555555555555555555555				2222 1111
555555555555555555555555555555555555555			4444 3333	2222 1111
555555555555555555555555555555555555555		**	4444 333	
*55555555555555555555555555555555555555			44444 33	
555555555555555555555555555555555555555			44444 33	
555555555555555555555555555555555555555			44444 33	
555555555555555555555555555555555555555				33 22222 1
555555555555555555555555555555555555555				33 22222 1
555555555555555555555555555555555555555				333 22222 1
*55555555555555555555555555555555555555				333 * 2222 *
				,,,,,

MAP LEGEND							
HORIZONTAL GRID NODE RANGE, FROM 1 TO 1	L						
VERTICAL GRID NODE RANGE, FROM 1 TO 11							
DEPENDENT VARIABLE RANGE MAP CHARACTER							

0.000E-01 -	1.000E-01	
1.0008-01 -	2.000E-01	1
2.0008-01 -	3.000E-01	
3.0002-01 -	4.000E-01	2
4.0008-01 -	5.000E-01	
5.000E-01 -	6.000E-01	3
6.000E-01 -	7.0002-01	
7.0008-01 -	8.000E-01	4
8.000E-01 -	9.000E-01	
9.0008-01 -	1.000E+00	5

÷. ‡

0.000	0.200	E LOCATIONS 0.400 E LOCATIONS	0.600	0.800	1.00	1.20	1.40	1.60	1.80	2.00
0.000	AXIS IS 1 0.200	POSITIVE UP	• THE PAGE 0.600	0.800	1.00	1.20	1.40	1.60	1.80	2.00
0.000	0.200			INTERSTITIA						2.00

VERTICAL SLICES

J = 1

11 10 9 8 7 6	1 5.8050E-03 5.6043E-03 5.3289E-03 5.6099E-03 7.7525E-03 1.3214E-02	2 1.7689E-02 1.7088E-02 1.6128E-02 1.6607E-02 2.1933E-02 3.5479E-02	3 3.0581E-02 2.9465E-02 2.7400E-02 2.7078E-02 3.3082E-02 4.9678E-02	4 4.5596E-02 4.3659E-02 3.9668E-02 3.7094E-02 4.0965E-02 5.5950E-02	5 6.4559E-02 6.1105E-02 5.3692E-02 4.6992E-02 4.6238E-02 5.6113E-02	6 9.0775E-02 8.4037E-02 7.0286E-02 5.6947E-02 4.9597E-02 5.2287E-02	7 0.1312 0.1158 8.9582E-02 6.6094E-02 5.0885E-02 4.6079E-02	8 0.2036 0.1605 0.1087 7.0920E-02 4.8175E-02 3.7726E-02	9 0.3643 0.2155 0.1152 6.3208E-02 3.7551E-02 2.5927E-02	10 0.8247 0.2228 7.3718E-02 3.0376E-02 1.5336E-02 9.5482E-03
5	2.23958-02	5.6518E-02	7.5087E-02	8.0709E-02	7.6885E-02	6.6560E-02	5.2617E-02	3.77588-02	2.29788-02	7.7931E-03
4	3.9561E-02	8.6821E-02	0.1060	0.1089	0.1023	8.8744E-02	6.9581E-02	4.77238-02	2.6993E-02	8.6132E-03
3	8.7745E-02	0.1410	0.1461	0.1387	0.1271	0.1123	9.2878E-02	6.74298-02	3.88298-02	1.2105E-02
2	0.2840	0.2362	0.1963	0.1687	0.1487	0.1337	0.1178	9.53238-02	6.1982E-02	2.0557E-02
1	0.6489	0.4450	0.2414	0.1873	0.1625	0.1512	0.1434	0.1303	0.1006	4.15258-02
	J = 2									
	1	2	3	4	5	6	7	8	9	10
11	5.8050E-03	1.7689E-02	3.0581E-02	4.55968-02	6.4559E-02	9.07758-02	0.1312	0.2036	0.3643	0.8247
10	5.6043E-03	1.7088E-02	2.9465E-02	4.3659E-02	6.1105E-02	8.4037E-02	0.1158	0.1605	0.2155	0.2228
9	5.3289E-03	1.6128E-02	2.7400E-02	3.9668E-02	5.3692E-02	7.0286E-02	8.9582E-02	0.1087	0.1152	7.3718E-02
8	5.6099E-03	1.6607E-02	2.70788-02	3.7094E-02	4.6992E-02	5.6947E-02	6.6094E-02	7.0920E-02	6.3208E-02	3.0376E-02
	7.7525E-03 1.3214E-02	2.1933E-02 3.5479E-02	3.3082E-02 4.9678E-02	4.0965E-02 5.5950E-02	4.6238E-02 5.6113E-02	4.9597E-02 5.2287E-02	5.0885E-02 4.6079E-02	4.8175E-02 3.7726E-02	3.7551E-02 2.5927E-02	1.5336E-02 9.5482E-03
6	2.23958-02	5.65182-02	<b>4.90762-02</b> <b>7.50872-02</b>	8.07098-02	7.6885E-02	6.65608-02	4.00/92-02 5.2617E-02	3.77588-02	2.29786-02	7.79316-03
4	3.9561E-02	8.68218-02	0.1060	0.1089	0.1023	8.8744E-02	6.95816-02	4.77238-02	2.69938-02	8.61328-03
1	8.7745E-02	0.1410	0.1461	0.1387	0.1271	0.1123	9.28788-02	6.74298-02	3.8829E-02	1.21056-02
2	0.2840	0.2362	0.1963	0.1687	0.1487	0.1337	0.1178	9.5323E-02	6.19828-02	2.05578-02
ī	0.6489	0.4450	0.2414	0.1873	0.1625	0.1512	0.1434	0.1303	0.1006	4.1525E-02

#### Y-DIRECTION - INTERSTITIAL PORE VELOCITY BETWEEN Y(J) AND Y(J+1) ( M/S)

.

# VERTICAL SLICES

- 67

	- 1	•

	1	. 2.	- Sec. 3	4	5	6 <b>6</b> 1	. 7	. 8	9	10
- 11	2.2403E-13	-1.3447E-13	2.0185E-13	-2.2448E-13	1.7977E-13	-6.7495E-14	9.0115E-14	-6.7679E-14	5.6463E-14	-2.2599E-14
				2.0346E-13						
9	1.4020E-13	-3.7389E-13	- 3.0384E-13	-3.7394E-13	1.8687E-13	-1.8661E-13	2.5587E-13	-1.3886E-13	6.8915E-14	-2.2797E-13
8	-2.5928E-13			5.0911E-14					-9.3623E-14	9.1965E-14
7	0.0000	-6.1727E-14	2.4221E-13	-1.1776E-13	2.8459E-13	-4.3825E-13	3.1509E-13	0.0000	0.0000	-4.1871E-13
6	0.0000	2.21532-13	-2.8816E-13	3.4678E-13	-3.9586E-13	5.5855E-13	-2.3126E-13	1.0714E-13	0.0000	1.4191E-13
- 5	4.0746E-13	-3.2432E-13	5.5905E-13	-7.7731E-14	7.4526E-14	-1.4014E-13	2.5754E-13	-5.8103E-14	3.1430E-13	-1.9362E-13
- 4	-3.3200E-13	8.2898E-13	-4.9497E-13	1.6302E-13	-3.1832E-13	6.0991E-13	-1.4187E-13	0.0000	-1.1167E-13	3.4950E-13
3	0.0000	-8.3127E-13	1.6607E-13	-4.9611E-13	1.6343E-13	-9.5516E-13	4.5397E-13	-6.9164E-13	1.2058E-13	-3.1214E-13
2	-1.6631E-13	1.6630E-13	0.0000	0.0000	-4.9488E-13	8.1083E-13	3.1261E-13	2.9193E-13	-1.2999E-13	1.1000E-13
1	0.0000	-4.9892E-13	0.0000	-1.6612E-13	6.6133E-13	0.0000	4.7367E-13	0.0000	1.3479E-13	-3.5013E-13

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. .....

11 11 0.0000 10 -3.3963E-14 9 -2.2724E-14 8 0.0000 7 2.7577E-13 6 -1.8559E-13 5 4.6986E-14 4 -3.3404E-13 3 0.0000 2 -2.9655E-13 1 0.0000

Z-DIRECTION - INTERSTITIAL PORE VELOCITY BETWEEN Z(K) AND Z(K+1) ( M/S)

#### VERTICAL SLICES

#### J = 1

	1	2	3	4	5	6	7	8	9	10
10	5.7866E-03	5.9248E-03	6.4297E-03	7.4921E-03	9.4663E-03	1.3093E-02	2.0197E-02	3.6207E-02	8.0299E-02	0.2301
9	1.6924E-02	1.7337E-02	1.8738E-02	2.1623E-02	2.6865E-02	3.6005E-02	5.1998E-02	8.0921E-02	0.1354	0.2376

.

6 5.2533E-02 5.1521E-02 4,9879E-02 5.0235E-02 5.4842E-02 6.5113E-02 8.1361E-02 0.1022 0.1241 0.1420 5 7.7592E-02 7.2491E-02 6.2906E-02 5.5513E-02 5.4215E-02 6.0676E-02 7.4727E-02 9.3692E-02 0.1126 0.1263	3	0.1994	0.1516	9.88698-02	6.30298-02	4.27662-02	3.5583E-02	4.0303E-02	5.5857E-02	7.6984E-02	9.4281E-02
5 7.7592E-02 7.2491E-02 6.2906E-02 5.5513E-02 5.4215E-02 6.0676E-02 7.4727E-02 9.3692E-02 0.1126 0.1263	3	•••••		9.88698-02	6.30298-02		3.5583E-02	4.0303E-02	5.5857E-02	7.6984E-02	9.4281E-02
	4	0.1207	0.1052	8.0368E-02	6.0270E-02	4.9664E-02	4.9680E-02	6.0153E-02	7.83958-02	9.7907E-02	0.1118

	11
10	0.8329
9	0.3873
8	0.2401
1	0.1798
6	0.1497
5	0.1313
4	0.1168
3	0.1009
2	7.8480E-02
1	3.92228-02

- -

**J = 2** 

	1	2	3	4	5	6	7	8	9	10
10	5.7866E-03	5.9248E-03	6.4297E-03	7.4921E-03	9.4663E-03	1.3093E-02	2.01978-02	3.6207E-02	8.0299E-02	0.2301
9	1.6924E-02	1.7337E-02	1.8738E-02	2.1623E-02	2.6865E-02	3.6005E-02	5.19988-02	8.0921E-02	0.1354	0.2376
8	2.7306E-02	2.7862E-02	2.9744E-02	3.3654E-02	4.0699E-02	5.2475E-02	7.1243E-02	0.1000	0.1420	0.1963
7	3.7916E-02	3.8247E-02	3.9625E-02	4.31498-02	5.01988-02	6.2180E-02	8.0286E-02	0.1049	0.1345	0.1638
6	5.2533E-02	5.1521E-02	4.9879E-02	5.0235E-02	5.4842E-02	6.5113E-02	8.13618-02	0.1022	0.1241	0.1420
5	7.7592E-02	7.2491E-02	6.2906E-02	5.5513E-02	5.4215E-02	6.0676E-02	7.4727E-02	9.36928-02	0.1126	0.1263
- 4	0.1207	0.1052	8.03682-02	6.0270E-02	4.9664E-02	4.9680E-02	6.01538-02	7.8395E-02	9.7907E-02	0.1118
3	0.1994	0.1516	9.88698-02	6.3029E-02	4.2766E-02	3.5583E-02	4.0303E-02	5.5857E-02	7.6984E-02	9.4281E-02
2	0.3963	0.2001	0.1053	5.6562E-02	3.11888-02	2.0501E-02	2.0146E-02	2.96062-02	4.7945E-02	6.8290E-02
1	0.6455	0.2806	7.1863E-02	2.7406E-02	1.1437E-02	5.0341E-03	3.4014E-03	5.9634E-03	1.3839E-02	2.75962-02

11 10 0.8329 9 0.3873 8 0.2401 7 0.1798 6 0.1497

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1999 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 - 1994 -

5	0.1313
4	0,1168
3	0,1009
2	7.8480E-02
1	3.9222E-02

4

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	***** JOB COMPLETED *****		
LAST TIME PLANE	CALCULATED	10.	(5)
LAST TIME PLANE	INDEX	50	

303

7.'

8.2.2. Heat Transport with Variable Density and Variable Viscosity

The second example problem is based upon thermal injection of a hot fluid through a well followed by production for an equivalent time period. The cylindrical coordinate system is used. Injection is for 90 days, followed by 90 days of withdrawal. Temperature of the injected fluid is 60 °C; flow rate is 203 2/s for both injection and withdrawal. Initial temperature in the aquifer is 20 °C. The other parameters are:

Well radius, 1 m Region outer radius, 246 m Aquifer thickness, 30 m Permeability,  $1.02 \times 10^{-10} \text{ m}^2$ Porosity, 0.35 Fluid compressibility, 0. Pa<sup>-1</sup> Porous matrix compressibility, 0. Pa<sup>-1</sup> Fluid heat capacity, 4,182 J/kg-°C Porous-medium heat capacity, 840 J/kg-°C Fluid density at 20 °C, 1,000 kg/m<sup>3</sup> Porous-medium density of the solid, 2,650 kg/m<sup>3</sup> Thermal conductivity of the fluid, 0.6 W/m-°C Thermal conductivity of the porous medium, 3.5 W/m-°C Coefficient of thermal expansion of the fluid, 0.375 °C<sup>-1</sup> Fluid viscosity at 20 °C, 0.001 kg/m-s Longitudinal dispersivity, 4 m Transverse dispersivity, 1 m

The well flow rate is to be allocated by mobility explicitly in time. The upper and lower boundaries are impermeable and thermally insulated. The boundary condition at the outer radius is specified as hydrostatic pressure with a specified temperature of 20 °C. The boundary condition pressures and temperatures do not vary with time.

Construct a numerical model of this system, and observe the movement of the injected hot water and subsequent withdrawal, for a total simulation time

of 180 days. Automatic spacing of 26 nodes in the radial direction and uniform spacing of 11 nodes in the vertical direction are to be used. Use backwards-in-time differencing with a fixed time step of 3 days. The cross-derivative dispersion terms should be calculated explicitly. Print results at 30 day intervals including contour maps of temperature. Use a contour range of 20 to 60 °C, with a contour interval of 5 °C.

The data file that will run example 2 is given in table 8.7. Again, the comments that do not pertain to this problem have been eliminated for brevity.

Table 8.8 contains selections from the output file for this problem. Only key results are presented with highlights of the output described. The initial heat in the region includes both the enthalpy of the fluid and of the porous matrix. The effective ambient permeability at the well bore is defined element by element, thus, 10 values occur. The well-flow rate is calculated explicitly at the beginning of each time step. The temperature field at the end of the injection period shows the less-dense fluid rising up over the cooler, resident fluid. Some numerical overshoot or spatial oscillation is apparent, with temperatures up to 62.5 °C appearing. This gives a zone of plus signs on the contour map of temperature. At the value of time step selected, only two cycles for solution of the pressure and temperature equations were needed, which is the minimum necessary for the explicit calculation of the cross-derivative dispersive-flux terms. During the production part of the cycle, we can see the preferential withdrawal of the warmer water from the upper part of the region, caused by the enhanced mobility of the water at higher temperature. An additional zone, shown as minus signs on the contour maps, shows that the temperature is below the lower limit of 20 °C selected for these plots. Again, this is a spatial-oscillation effect, caused by the coarseness of the grid in conjunction with central differencing for the advective terms. The global-balance summary shows that, at the end of 90 days of withdrawal, about 86 percent of the heat has been recovered. Only about 0.1 percent of the heat left the region through the boundary at the exterior radius. The fluid-withdrawal and heat-withdrawal rates are shown on a perlayer basis at the end of the simulation. The well is producing water at about 38 °C at this time.

Table 8.7.--Input-data file for example problem 2

C....HST DATA-INPUT FORM C... NOTES: C... INPUT LINES ARE DENOTED BY C.N1.N2.N3 WHERE HI IS THE READ GROUP HUNBER, N2.N3 IS THE RECORD NUMBER C... A LETTER INDICATES AN EXCLUSIVE RECORD SELECTION MUST BE MADE C... C... I.E. A OR B OR C (0) - OPTIONAL DATA WITH CONDITIONS FOR REQUIREMENT C... A RECORD NUMBER IN SQUARE BRACKETS IS THE RECORD WHERE THAT PARAMETER IS FIRST SET C... C..... INPUT BY I.J.K RANGE FORMAT IS: C.O.1.. 11,12,J1,J2,K1,K2 C.O.2.. VAR1, INOD1, [VAR2, INOD2, VAR3, INOD3] C... USE AS MANY OF LINE 0.1 & 0.2 SETS AS NECESSARY C ... END WITH LINE 0.3 C.O.3.. 0 / THE SPACE IS REQUIRED C... (NNN) - INDICATES THAT THE DEFAULT NUMBER, NNN, IS USED IF A ZERO IS ENTERED FOR THAT VARIABLE C... (T/F) - INDICATES A LOGICAL VARIABLE C... [1] - INDICATES AN INTEGER VARIABLE C.....START OF THE DATA FILE C.... DIMENSIONING DATA - READI C.1.1 .. TITLE LINE 1 EXAMPLE #2 THERMAL INJECTION AND WITHDRAWAL. CYLINDRICAL SYSTEM. C.1.2 .. TITLE LINE 2 VARIABLE DENSITY AND VISCOSITY C.1.3 .. RESTRT(T/F), TIMRST F. / C.1.4 .. HEAT.SOLUTE.EEUNIT.CYLIND.SCALMF: ALL (T/F) TEETE C.1.5 .. NX,NY,NZ,NHCN 26 1 11 0 C.1.6 .. NPTCBC\_NFBC\_NAIFC\_NLBC\_NHCBC\_NHEL 11 0 0 0 0 1 C.1.7 .. NPHZ C.1.8 .. SLMETH[1],LCROSD(T/F) 1 F C.1.9 .. IBC BY I.J.K RANGE (0.1-0.3) .WITH NO INOD PARAMETER. FOR EXCLUDED CELLS 0/ C.1.10 .. RDECHO(T/F) C.....STATIC DATA - READ2 C....OUTPUT INFORMATION C.2.1 .. PRTRE(T/F) F

C....COORDINATE GEOMETRY INFORMATION C.....CYLINDRICAL COORDINATES C.2.2B.1A .. R(1),R(MR),ARGRID(T/F);(0) - CYLIND [1.4] 1. 246. T C.2.28.18 .. R(1);(0) - NOT ARGRID [2.28.1A];(0) - CYLIND [1.4] C.2.28.2 .. UNIGRZ(T/F);(0) - CYLIND [1.4] T' C.2.2B.3A .. Z(1), Z(NZ); (0) - UNIGRZ [2.2B.3A], CYLIND [1.4] 0. 30. C.2.2B.3B .. Z(K);(0) - NOT UNIGRZ [2.2B.3A],CYLIND [1.4] C.....FLUID PROPERTY INFORMATION C.2.4.1 .. BP 0. C.2.4.2 .. PO,TO,MO,DENFO 0. 20. 0. 1000. C.2.5.1 .. NOTVO, TVFO(I), VISTFO(I), I=1 TO NOTVO; (0) - HEAT [1.4] OR HEAT [1.4] AND SOLUTE [1.4] OR .NOT.HEAT AND .NOT.SOLUTE [1.4] 2 20. .001 60. 4.62E-4 C.....REFERENCE CONDITION INFORMATION C.2.6.1 .. PAATH 0. C.2.6.2 .. POH, TOH 0. 20. C.....FLUID THERMAL PROPERTY INFORMATION C.2.7 .. CPF\_KTHF\_BT;(0) - HEAT [1.4] 4182. .6 3.75E-4 C.....POROUS MEDIA ZONE INFORMATION C.2.9.1 .. IPMZ\_I1Z(IPMZ).IZZ(IPMZ).J1Z(IPMZ).J2Z(IPMZ).K1Z(IPMZ).K2Z(IPMZ) 1 1 26 1 1 1 11 C..... USE AS MANY 2.9.1 LINES AS NECESSARY C.2.9.2 .. END WITH 0 / 0/ C.....POROUS MEDIA PROPERTY INFORMATION C.2.10.1 .. KXX(IPHZ),KYY(IPHZ),KZZ(IPHZ),IPHZ=1 TO NPHZ [1.7] 1.02E-10,,1.02E-10 C.2.10.2 .. POROS(IPMZ), IPMZ=1 TO NPMZ [1.7] .35 C.2.10.3 .. ABPH(IPMZ), IPMZ=1 TO NPMZ [1.7] 0. · C.....POROUS MEDIA THERMAL PROPERTY INFORMATION C.2.11.1 .. RCPPH(IPMZ), IPMZ=1 TO NPHZ [1.7]:(0) - HEAT [1.4] 2.22686 C.2.11.2 .. KTXPH(IPMZ),KTYPH(IPMZ),KTZPH(IPMZ),IPMZ=1 TO NPMZ [1.7];(0) - HEAT [1.4] 3\*3.5 C.....POROUS MEDIA SOLUTE AND THERMAL DISPERSION INFORMATION C.2.12 .. ALPHL(IPMZ), ALPHT(IPMZ), 1PMZ=1 TO NPMZ [1.7]; (0) - SOLUTE [1.4] OR HEAT [1.4]

```
4. 1.
C.....SOURCE-SINK WELL INFORMATION
C.2.14.1 .. RDWDEF(T/F);(0) - NWEL [1.6] > 0
C.2.14.3. .. IWEL, IW, JW, LCBOTW, LCTOPW, WBOD, WOMETH[I]; (0) - RDWDEF [2.14.1],
1 1 1 1 11 2. 11
C.2.14.4 .. WCF(L);L = 1 TO NZ (EXCLUSIVE) BY ELEMENT
10*1.
C.....USE AS MANY 2.14.3-6 LINES AS NECESSARY
C.2.14.7 .. END WITH 0 /
0/
C....BOUNDARY CONDITION INFORMATION
           SPECIFIED VALUE B.C.
C....
C.2.15 .. IBC BY I.J.K RANGE (0.1-0.3) WITH NO INOD PARAMETER; (0) - NPTCBC [1.6] > 0
26 26 1 1 1 11
100
0/
C....FREE SURFACE B.C.
C.2.20 .. FRESUR(T/F), PRTCCH(T/F)
·F /
C....INITIAL CONDITION INFORMATION
C.2.21.1 .. ICHYOP, ICT, ICC; ALL (T/F); IF NOT. HEAT, ICT = F, IF NOT. SOLUTE, ICC = F
TTF
C.2.21.3A .. ZPINIT, PINIT; (0) - ICHYDP [2.21.1] AND NOT ICHWT [2.21.2]
30. 0.
C.2.21.48 .. T BY I, J, K RANGE (0.1-0.3); (0) - HEAT (1.4) AND ICT (2.21.1)
1 26 1 1 1 11
20. 1
0/
C....CALCULATION INFORMATION
C.2.22.1 .. FDSMTH_FDTMTH
.5 1.
C.2.22.2 .. TOLDEN(.001), MAXITN(5)
0. 10
C....OUTPUT INFORMATION
C.2.23.1 .. PRTPMP, PRTFP, PRTIC, PRTBC, PRTSLM, PRTWEL; ALL (T/F)
6*T
C.2.23.2... IPRPTC, PRTDV(T/F);(0) - PRTIC [2.23.1]
110 F
C.2.23.4 .. PLTZON(T/F);(0) - PRTPHP [2.23.1]
C.2.23.5 .. OCPLOT(T/F)
C..... TRANSIENT DATA - READ3
```

\*\*\*

1

C.3.1 .. THRU(T/F) C....IF THRU IS TRUE PROCEED TO RECORD 3.99 C....THE FOLLOWING IS FOR NOT THRU C.....SOURCE-SINK WELL INFORMATION C.3.2.1 .. RDWFLO(T/F).RDMHD(T/F);(0) - NWEL [1.6] > 0 T F C.3.2.2 .. INEL, OWV, PWSUR, PWKT, TWSRKT, CWKT; (0) - RDWFL0 [3.2.1] OR RDWHD [3.2.1] 1 203. 0. 0. 60. 0. C..... USE AS MANY 3.2.2 LINES AS NECESSARY C.3.2.3 .. END WITH 0 / 0/ C....BOUNDARY CONDITION INFORMATION SPECIFIED VALUE B.C. C.... C.3.3.1 .. RDSPBC, RDSTBC, RDSCBC, ALL(T/F); (0) ~ NOT CYLIND [1.4] AND NPTCBC [1.6] > 0 TFF C.3.3.2 .. PMP B.C. BY I.J.K RANGE (0.1-0.3);(0) - RDSPBC [3.3.1] 26 26 1 1 11 11 0.1 26 26 1 1 10 10 2.9421E4 1 26 26 1 1 9 9 5.8842E4 1 26 26 1 1 8 8 8.826364 1 26 26 1 1 7 7 1.17684E5 1 26 26 1 1 6 6 1.47105E5 1 26 26 1 1 5 5 1.7652685 1 26 26 1 1 4 4 2.05947E5 1 26 26 1 1 3 3 2.35368E5 1 26 26 1 1 2 2 2.6478985 1 26 26 1 1 1 1 2.942165 1 0/ C.3.3.3 .. TSBC BY I, J, K RANGE (0.1-0.3); (0) - RDSPBC [3.3.1] AND HEAT [1.4] 26 26 1 1 1 11 20.1 0/ C....CALCULATION INFORMATION

```
C.3.7.1 .. RDCALC(T/F)
C.3.7.2 .. AUTOTS(T/F);(0) - ROCALC [3.7.1]
C.3.7.3.A .. DELTIN; (0) - RDCALC [3.7.1] AND NOT AUTOTS [3.7.2]
2.59265
C.3.7.4 .. TINCHG
7.776086
C....OUTPUT INFORMATION
C.3.8.1 .. PRIVEL, PRIOV, PRISLN, PRIKD, PRIPTC, PRIGFB, PRIWEL, PRIBCF; ALL [1]
0 0 10 0 10 -1 0 30
C.3.8.2 .. IPRPTC;(0) - IF PRIPTC [3.8.1] NOT = 0
010
C.3.8.3 .. CHKPTD(T/F), NTSCHK, SAVLDO(T/F)
F /
C....CONTOUR HAP INFORMATION
C.3.9.1 .. RDNPDT, PRTNPD; ALL (T/F)
TT
C.3.9.2 .. NAPPTC_PRINAP[1];(0) - RDNPDT [3.9.1]
010 10
C.3.9.3 .. YPOSUP(T/F), ZPOSUP(T/F), LENAX, LENAY, LENAZ; (0) - ROMPOT [3.9.1]
FT 12. 0. 6.
C.3.9.4 .. IMAP1(1), IMAP2(NX), JMAP1(1), JMAP2(NY), KMAP1(1), KMAP2(NZ), AMIN, AMAX, NMPZON(5); (0) - RDMPDT [3.9.1]
00000020.60.8
C....ONE OF THE 3.9.4 LINES REQUIRED FOR EACH DEPENDENT VARIABLE
C....
           TO BE NAPPED
C....END OF FIRST SET OF TRANSIENT INFORMATION
C. .
C.....READ SETS OF READ3 DATA AT EACH TINCHG UNTIL THRU (LINES 3.N1.N2)
C..... TRANSIENT DATA - READ3
C.3.1 .. THRU(T/F)
C....IF THRU IS TRUE PROCEED TO RECORD 3.99
C....THE FOLLOWING IS FOR NOT THRU
C.....SOURCE-SINK HELL INFORMATION
C.3.2.1 .. RDWFLO(T/F), RDWHD(T/F); (0) - NWEL [1.6] > 0
T F
C.3.2.2 .. IVEL.QUV.PWSUR.PWKT.TWSRKT.CWKT;(0) - RDWFL0 [3.2.1] OR RDWH0 [3.2.1]
1 -203. 0. 0. 60. 0.
C..... USE AS MANY 3.2.2 LINES AS NECESSARY
C.3.2.3 .. END WITH 0 /
0/
C....BOUNDARY CONDITION INFORMATION
C....
           SPECIFIED VALUE B.C.
C.3.3.1 .. RDSPBC, RDSTBC, RDSCBC, ALL(T/F); (0) - NOT CYLIND [1.4] AND NPTCBC [1.6] > 0
```

```
FFF
C....CALCULATION INFORMATION
C.3.7.1 .. RDCALC(T/F)
C.3.7.2 .. AUTOTS(T/F);(0) - ROCALC [3.7.1]
C.3.7.3.A .. DELTIM; (0) - ROCALC [3.7.1] AND NOT AUTOTS [3.7.2]
2.59215
C.3.7.4 .. TINCHG
1.5552E7
C....OUTPUT INFORMATION
C.3.8.1 .. PRIVEL, PRIDY, PRISLN, PRIKD, PRIPTC, PRIGFB, PRIWEL, PRIBCF; ALL [1]
0 0 10 0 10 -1 -1 30
                                                                .
C.3.8.2 .. IPRPTC; (0) - IF PRIPTC [3.8.1] NOT = 0
010
C.3.8.3 .. CHKPTD(T/F), NTSCHK, SAVLDO(T/F)
F /
C....CONTOUR MAP INFORMATION
C.3.9.1 .. ROMPDT, PRTNPD; ALL (T/F)
TT
C.3.9.2 .. MAPPTC, PRIMAP[1];(0) - RDMPDT [3.9.1]
010 10
C.3.9.3 .. YPOSUP(T/F), ZPOSUP(T/F), LENAX, LENAY, LENAZ; (0) - ROMPOT [3.9.1]
FT 12. 0. 6.
C.3.9.4 .. IMAP1(1), IMAP2(NX), JMAP1(1), JMAP2(NY), KMAP1(1), KMAP2(NZ), AMIN, AMAX, NMPZON(5); (0) - RDMPDT (3.9.1)
0 0 0 0 0 0 20. 60. 8
C....ONE OF THE 3.9.4 LINES REQUIRED FOR EACH DEPENDENT VARIABLE
C..... TO BE MAPPED
C....END OF SECOND SET OF TRANSIENT INFORMATION
C.....READ SETS OF READ3 DATA AT EACH TIMCHG UNTIL THRU (LINES 3.N1.N2)
C....END OF CALCULATION LINES FOLLOW, THRU-, TRUE.
C.3.99.1 .. THRU
C.....TEMPORAL PLOT INFORMATION
C.3.99.2 .. PLOTWP, PLOTWT, PLOTWC; ALL (T/F)
3*F
C....END OF DATA FILE
```

Table 8.8.--Selections from the output file for example problem 2

\* THREE DINENSIONAL FLOW, HEAT AND SOLUTE • TRANSPORT SINULATOR - (HST3D):RELEASE - 1.0

EXAMPLE #2 THERMAL INJECTION AND WITHDRAWAL, CYLINDRICAL SYSTEM. VARIABLE DENSITY AND VISCOSITY

\*\*\* FUNDAMENTAL INFORMATION \*\*\* CYLINDRICAL COORDINATES HEAT TRANSPORT SIMULATION NO SOLUTE TRANSPORT SIMULATON WELLBORE MODEL MAY INCLUDE : FLUID PRESSURE DROP HEAT BALANCE UP THE RISER PIPE INPUT DATA IS EXPECTED IN METRIC UNITS

**\*\*\*** PROBLEM DIMENSIONING INFORMATION **\*\*\*** 26 1 11 NUMBER OF POROUS MEDIA ZONES ..... NPMZ . 1 NUMBER OF SPECIFIED PRESSURE, TEMPERATURE OR MASS FRACTION B.C. ...... NPTCBC 11 NUMBER OF SPECIFIED FLUX B.C. CELLS (FLOW, HEAT OR SOLUTE)...... NFBC . 0 NUMBER OF HEAT CONDUCTION B.C. CELLS ...... NHCBC Δ NUMBER OF NODES OUTSIDE REGION FOR EACH HEAT CONDUCTION B.C. CELL ...... NHCN A 0 0 1

#### DIRECT D4 SOLVER IS SELECTED FULL CROSS-DISPERSIVITY COEFFICIENT STORAGE ALLOCATED THE A4 ARRAY IN DADES IS DIMENSIONED ..... 2917 ELEMENTS THE TOTAL STORAGE REQUIRED BY THE DIRECT METHOD IS ..... 4347 ELEMENTS THE TOTAL STORAGE REQUIRED BY THE ITERATIVE METHOD IS ..... 2002 ELEMENTS TOTAL LENGTH OF LABELED COMMON BLOCKS ..... 6681 BYTES REQUIRED COMPILED LENGTH OF VARIABLE LENGTH REAL ARRAY (VPA ARRAY) ..... 13876 ELEMENTS 250000 ELEMENTS LENGTH OF VARIABLE LENGTH INTEGER ARRAY (IVPA ARRAY) ..... 2099 ELEMENTS 20000 ELEMENTS

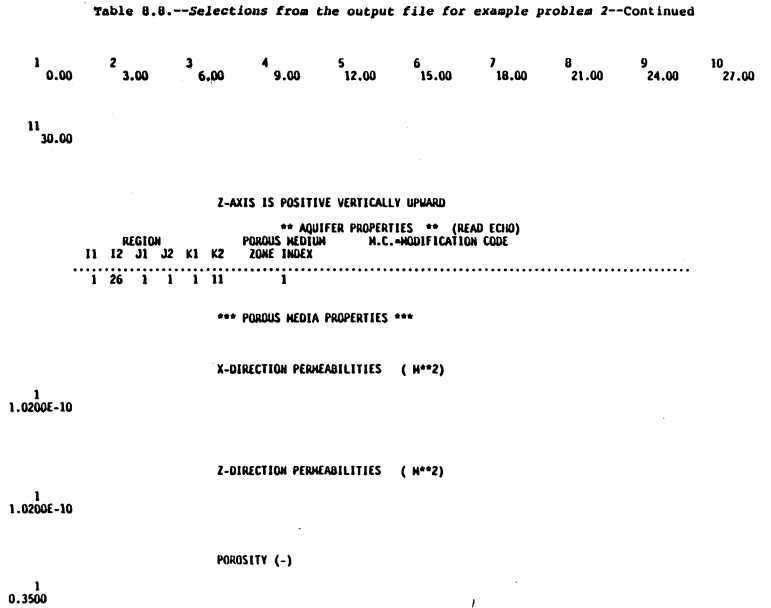
# Table 8.8.--Selections from the output file for example problem 2--Continued

	AI AI	, <del>***</del>	CYLINDRICAL	ANT OR STATIO (R-Z) COORDI	INATE DATA **	1.000 ( 246.0 (	M) M)		
·		R-D	IRECTION NOD	E COORDINATES	5 (M)				
1 1.00	2 1.25	3 1.55	4 1.94	5 2.41	6 3.01	7 3.75	8 4.67	9 5.82	10 7.26
11 9.04	12 11.27	13 14.05	14 17.51	15 21.82	16 27.20	17 33.90	18 42.25	19 52.66	20 65.63
21 81.80	22 101.95	23 127.06	24 158.37	25 197.38	26 246.00				
		R-CI	DORDINATE CEI	LL BOUNDARY (	LOCATIONS (BI	ETWEEN NODE(1	I) AND NODE(I	(+1)) ( M)	
1 1.12	2 1.39	3 1.74	4 2.17	5 2.70	6 3.36	7 4.19	8 5.23	9 6.51	10 8.12
11 10.12	12 12.61	13 15.72	14 19.59	15 24.41	16 30.43	17 37.92	18 47.26	19 58.91	20 73.42
21 91.50	22 114.05	23 142.14	24 177.16	25 220.80		· :	<b>-</b> .		

1

Z-DIRECTION NODE COORDINATES ( M)

313



(

#### Table 8.8.--Selections from the output file for example problem 2--Continued

#### \*\*\* INTERHEDIATE COMPUTED DATA \*\*\*

#### R-DIRECTION CONDUCTANCE FACTOR BETWEEN R(I) AND R(I+1) (M\*\*3)

VERTICAL SLICES

315

J = 1

	1.0	2	<b>3</b> - 19 - 19	• • • • <b>4</b>	° 5 °	6	7	8	9	10
11	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09
10	8.7309E-09	8.7309E-09	8,7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09
9	- 8.7309E-09		8.7309E-09		8.7309E-09					8.7309E-09
8	8.7309E-09	8,7309E-09	8.73098-09	8.7309E-09	8.7309E-09	8.73092-09	8.7309E-09	8.7309E-09	8.7309E-09	8.73032-09
7	8.7309E-09	8.7309E-09	8,73095-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8,7309E-09	8.7309E-09	8.7309E-09
6	8.7309E-09	8.7309E-09	8.7309E-09	8.73095-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09
5	8.7309E-09	8,73092-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09
° 4	8.7309E-09	8.7309E-09	8.7309E-09	8.73098-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09
3	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.73098-09	8.73092-09	8.7309E-09	8.7309E-09	8.7309E-09
2	8.7309E-09	8,7309E-09	8.7309E-09	8.7309E-09	8.73092-09	8.7309E-09	8.73092-09	8.7309E-09	8.7309E-09	8.7309E-09
1	4.3654E-09	4,3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09

12 15 11 13 14 16 17 18 19 20 11 4,3654E-09 9 8.7309E-09 8 8.7309E-09 7 8.7309E-09 6 8.7309E-09 8.73098-09 8.7309E-09 8.7309E-09 8.7309E-09 8.7309E-09 8.7309E-09 8.7309E-09 5 8.7309E-09 4 8.7309E-09 8.7309E-09 8.7309E-09 8.7309E-09 3 8.7309E-09 2 8.7309E-09 1 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09 4.3654E-09

	21	22	23	24	25
11	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09	4.3654E-09
10	8.7309E-09	8.7309E-09	8.7309E-09	8.73092-09	8.7309E-09
9	8.7309E-09	8.73098-09	8.7309E-09	8.7309E-09	8.7309E-09
8	8.7309E-09	8.73092-09	8.7309E-09	8.7309E-09	8.7309E-09

Table 8.8.--Selections from the output file for example problem 2--Continued

7	8.7309E-09	8.73092-09	8.73098-09	8.73092-09	8.73092-09
6	8.7309E-09	8.73092-09	8.73092-09	8.7309E-09	8.73092-09
5	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.73092-09
4	8.7309E-09	8.7309E-09	8.7309E-09	8.73092-09	8.7309E-09
3	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09	8.7309E-09
2	8.7309E-09	8.73098-09	8.7309E-09	8.73098-09	8.7309E-09
1	4.36548-09	4.36548-09	4.36548-09	4.3654E-09	4.3654E-09

#### Z-DIRECTION CONDUCTANCE FACTOR BETWEEN Z(K) AND Z(K+1) ( N++3)

.

VERTICAL SLICES

J = 1

.

	1	2	3	4	5	6	. 7	8	9	10
10	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.68988-10	1.0392E-09	1.6142E-09	2.5075E-09
9	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
8	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
7	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
6	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
5	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
- 4	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
3	2.6852E-11	7.3967E-11	1.14908-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
2	2.6852E-11	7.3967E-11	1.1490E-10	1.7848E-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
1	2.6852E-11	7.3967E-11	1.14902-10	1.78482-10	2.7724E-10	4.3066E-10	6.6898E-10	1.0392E-09	1.6142E-09	2.5075E-09
	11	12	13	14	15 2 26705 00	16	17	18	19	20
10	3.8950E-09 3.8950E-09	6.0504E-09 6.0504E-09	9.3985E-09 9.3985E-09	1.4599E-08 1.4599E-08	2.2678E-08 2.2678E-08	3.5228E-08 3.5228E-08	5.4721E-08 5.4721E-08	8.5003E-08 8.5003E-08	1.3204E-07 1.3204E-07	2.0511E-07 2.0511E-07
9 8	3.89502-09	6.0504E-09	9.39852-09	1.45992-08	2.26782-08	3.5228E-08	5.47216-08	8.50032-08	1.32042-07	2.05112-07
07	3.89502-09	6.0504E-09	9.39858-09	1.45998-08	2.26782-08	3.5228E-08	5.47212-08	8.50032-08	1.32042-07	2.05112-07
6	3.89502-09	6.05042-09	9.39858-09	1.45998-08	2.26782-08	3.52288-08	5.47216-08	8.50032-08	1.3204E-07	2.0511E-07
5	3.89502-09	6.05042-09	9.3985E-09	1.45998-08	2.26788-08	3.52288-08	5.47218-08	8.5003E-08	1.3204E-07	2.05116-07
4	3.8950E-09	6.0504E-09	9.39858-09	1.45998-08	2.2678E-08	3.5228E-08	5.47216-08	8.50032-08	1.3204E-07	2.0511E-07
3	3.8950E-09	6.0504E-09	9.3985E-09	1.45998-08	2.2678E-08	3.52282-08	5.47212-08	8.50032-08	1.32042-07	2.0511E-07
2	3.8950E-09	6.0504E-09	9.39858-09	1.45998-08	2.26782-08	3.52282-08	5.4721E-08	8.5003E-08	1.32048-07	2.05116-07
ī	3.89502-09	6.0504E-09	9.39858-09	1.4599E-08	2.2678E-08	3.5228E-08	5.4721E-08	8.5003E-08	1.32048-07	2.0511E-07

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10	3.1861E-07	4.9492E-07	7.6879E-07	1.1942E-06	1.8551E-06	1.2566E-06
9	3.1861E-07	4.9492E-07	7.6879E-07	1.1942E-06	1.8551E-06	1.2566E-06
8	3.1861E-07	4.9492E-07	7.6879E-07	1.1942E-06	1.8551E-06	1.2566E-06
7	3.1861E-07	4.94928-07	7.6879E+07	1.1942E-06	1.8551E-06	1.2566E-06
6	3.1861E-07	4.9492E-07	7.68792-07	1.1942E-06	1.8551E-06	1.2566E-06
5	3.1861E-07	4.9492E-07	7.6879E-07	1.19428-06	1.8551E-06	1.2566E-06
4	3.1861E-07	4.9492E-07	7.68792-07	1.1942E-06	1.8551E-06	1.2566E-06
3	3.1861E-07	4.9492E-07	7.6879E-07	1.1942E-06	1.8551E-06	1.2566E-06
2	3.1861E-07	4.9492E-07	7.6879E-07	1.1942E-06	1.8551E-06	1.2566E-06
1	3.1861E-07	4.9492E-07	7.6879E-07	1.1942E-06	1.8551E-06	1.2566E-06

\*\*\* PROPERTIES BY POROUS MEDIUM ZONE \*\*\*

POROUS MEDIUM VERTICAL COMPRESSIBILITY (1/ PA)

0.0000

DENSITY-HEAT CAPACITY PRODUCT ( J/ M\*\*3-DEG.C)

1 2.2260E+06

THERMAL CONDUCTIVITY IN X-DIRECTION ( W/ M-DEG.C)

1 3.500

> 1 3.500

> > THERMAL CONDUCTIVITY IN Z-DIRECTION ( W/ M-DEG.C)

1 3.500

LONGITUDINAL DISPERSIVITY ( M)

1 4.000

## TRANSVERSE DISPERSIVITY ( H)

1 1.000

ATHOSPHERIC PRESSURE (ABSOLUTE)	PAATH	101325.0	( PA)
REFERENCE PRESSURE FOR ENTHALPY	POH	0.0	( PA)
REFERENCE TEMPERATURE FOR ENTHALPY	TOH	20.0	(DEG.C)

+++ FLUID PROPERTIES +++ PHYSICAL FLUID COMPRESSIBILITY	BP	0.008-01	(1/ PA)
REFERENCE PRESSURE FOR DENSITY REFERENCE TEMPERATURE FOR DENSITY FLUID DENSITY AT SOLUTE NASS FRACTION OF ZERO	TO	0.0 20.0 1000.	( PA) (DEG.C) (KG/ H**3)
THERNAL FLUID COEFFICIENT OF THERNAL EXPANSION FLUID HEAT CAPACITY FLUID THERNAL CONDUCTIVITY FLUID SPECIFIC ENTHALPY AT REFERENCE CONDITIONS	CPF KTHF .	4.182E+03 0.600	(1/DEG.C) ( J/KG-DEG.C) ( W/ M-DEG.C) ( J/KG)

VISCOSITY-TEMPERATURE DATA TABLE TEMPERATURE (DEG.C) VISCOSITY (KG/ H-S)

FLUID AT SOLUTE WINIHUM MASS FRACTION

 20.0		1.0008-03
60.0	·	4.6208-04

\*\*\* INITIAL CONDITIONS \*\*\*

INITIAL AQUIFER FLUID PRESSURE FOR HYDROSTATIC I.C PINIT.	0.0 (	( PA)
ELEVATION OF INITIAL PRESSURE ZPINIT	30.0 (	(H)

INITIAL PRESSURE DISTRIBUTION ( PA)

### VERTICAL SLICES

#### J = 1

	1	2	3	4	5	6 <sup>`</sup>	·7	8	9	10
11	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
10	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04
9	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04
8	8.8263E+04	8,8263E+04	8.8263E+04	8.8263E+04	8,8263E+04	8.8263E+04	8.8263E+04	8.82638+04	8.8263E+04	8.8263E+04
7	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05	1.1768E+05
6	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05	1.4711E+05
- 5	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05	1.7653E+05
- 4	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05	2.0595E+05
3	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05	2.3537E+05
2	2.6479E+05	2.6479E+05	2.6479E+05	2.6479E+05	2.6479E+05	2.6479E+05	2.64795+05	2.6479E+05	2.6479E+05	2.6479E+05
1	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05	2.9421E+05

12 14 15 16 17 18 19 20 11 13 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 11 0.0000 0.0000 0.0000 10 2.9421E+04 9 5.8842E+04 8 8.8263E+04 1.1768E+05 1.1768E+05 7 1.1768E+05 1.1768E+05 1.1768E+05 1.1768E+05 1.1768E+05 1.1768E+05 1.1768E+05 1.1768E+05 1.4711E+05 1.4711E+05 1.4711E+05 1.4711E+05 1.4711E+05 1.4711E+05 1.4711E+05 1.4711E+05 6 1.4711E+05 1.4711E+05 5 1.7653E+05 4 2.0595E+05 .3 2.3537E+05 2 2.6479E+05 1 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05 2.9421E+05

	21	22	23	24	25	26
11	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
10	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04	2.9421E+04
9	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04	5.8842E+04
8	8.8263E+04	8.8263E+04	8.8263E+04	8.8263E+04	8.8263E+04	8.8263E+04

68E+05
11E+05
53E+05
95E+05
37E+05
79E+05
21E+05

INITIAL TEMPERATURES (DEG.C)

# VERTICAL SLICES

J = 1

	1	2	3	4	5	6	7	8	9	10
11	- 20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
10	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
ğ	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
9 8	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
ž	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
6	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
6 5	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
Ă	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
3	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
3	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
ī	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
•										
	11	12	13	14	15	16	17	18	19	20
11	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
10	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
10 9	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
8	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
7	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
6	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
5	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
4	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
3	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
2	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0
1	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0	20.0

<u>, 7</u>

	21	22	23	24	25	26
11	20.0	20.0	20.0	20.0	20.0	20.0
10	20.0	20.0	20.0	20.0	20.0	20.0
9	20.0	20.0	20.0	20.0	20.0	20.0
8	20.0	20.0	20.0	20.0	20.0	20.0
7	20.0	20.0	20.0	20.0	20.0	20.0
6	20.0	20.0	20.0	20.0	20.0	20.0
5	20.0	20.0	20.0	20.0	20.0	20.0
4	20.0	20.0	20.0	20.0	20.0	20.0
3	20.0	20.0	20.0	20.0	20.0	20.0
2	20.0	20.0	20.0	20.0	20.0	20.0
ī	20.0	20.0	20.0	20.0	20.0	20.0

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# INITIAL PORE VOLUME PER CELL RING ( M\*\*3)

VERTICAL SLICES

•	• ·	•				j =	1				
		1	2	3	4	5	6	7	8	9	10
	11	0.4146	1.142	1.774	2.756	4.281	6.650	10.33	16.05	24.93	38.72
11	10	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	9	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	8	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	7	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	6	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	- 5	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	4	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	3	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	2	0.8292	2.284	3.548	5.512	8.562	13.30	20.66	32.09	49.85	77.44
	1	0.4146	1.142	1.774	2.756	4.281	6.650	10.33	16.05	24.93	38.72
	•	11	12	13	14	15	16	17	18	19	20
	11	60.14	93.43	145.1	225.4	350.2	544.0	845.0	1313.	2039.	3167.
	10	120.3	186.9	290.2	450.9	700.4	1088.	1690.	2625.	4078.	6334.
	9	120.3	186.9	290.2	450.9	700.4	1088.	1690.	2625.	4078.	6334.
	8	120.3	186.9	290.2	450.9	700.4	1088.	1690.	2625.	4078.	6334.
	7	120.3	186.9	290.2	450.9	700.4	1088.	1690.	2625.	4078.	6334.
	6	120.3	186.9	290.2	450.9	700.4	1088.	1690.	2625.	4078.	6334.
	5	120.3	186.9	290.2	450.9	700.4	1088.	1690.	2625.	4ù78.	6334.

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4	120.3 120.3	186.9 186.9	290.2 290.2	450.9 450.9	700.4 700.4	1088. 1088.	1690. 1690.	2625. 2625.	4078. 4078.	6334. 6334.
2 1	120.3 60.14	186.9 93.43	290.2 145.1	450.9 225.4	700.4 350.2	1088. 544.0	1690. 845.0	2625. 1313.	4078. 2039.	6334. 3167.
	21	22	23	24	25	26				
- 11	4920.	7642.	1.1871E+04	1.8440E+04	2.8644E+04	1.9404E+04				
10	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
ģ	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
8	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
7	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
6	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.72882+04	3.8808E+04				
5	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
4	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
3	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.72888+04	3.8808E+04				
2	9839.	1.5284E+04	2.3742E+04	3.6880E+04	5.7288E+04	3.8808E+04				
ī	4920.	7642.	1.1871E+04	1.8440E+04	2.8644E+04	1.9404E+04				

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#### \*\*\* WELL DATA \*\*\* .... PERFORATIONS CALCULATION WELL DIAMETER WELL TYPE J K1 K2 (H) NO. I 2.0008+00 1 1 1 1 11 11 WELL NO. 1 SPECIFIED FLOW RATE ALLOCATION BY NOBILITY ELENENT LEVEL EFFECTIVE AMBIENT LAYER **WELL FLOW FACTOR** PERNEABILITY NO. NO. ( H\*\*2) ( H\*\*3) WELL NO. 1 1.530E-10 3.060E-10 11 1.0208-10 10 10 1.020E-10 9 9 3.060E-10

......

8	1.0201-10	8	3.0601-10
7	1.020E-10	7	3.060E-10
6	1.020E-10	6	3.060E-10
5	1.020E-10	5	3.060E-10
4	1.020E-10	4	3.060E-10
3	1.020E-10	3	3.060E-10
2	1.020E-10	2	3.060E-10
1	1.020E-10	· 1	1.530E-10

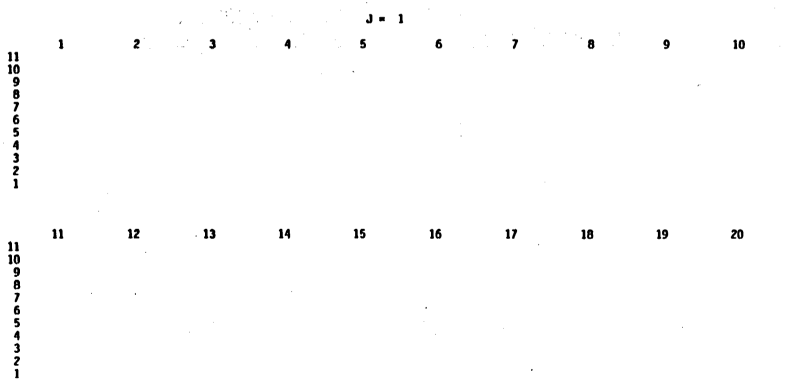
EXPLICIT WELL FLOW RATE AT EACH LAYER

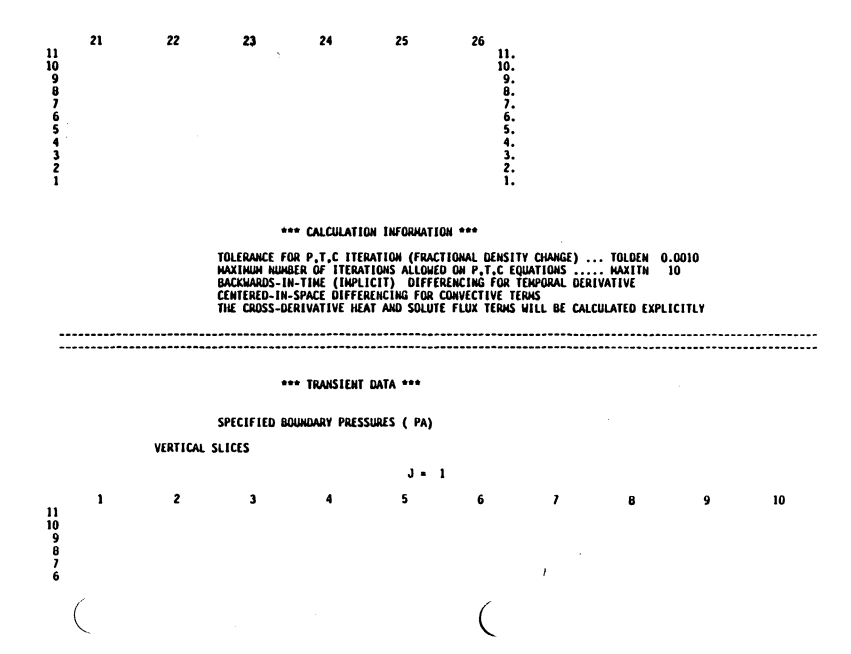
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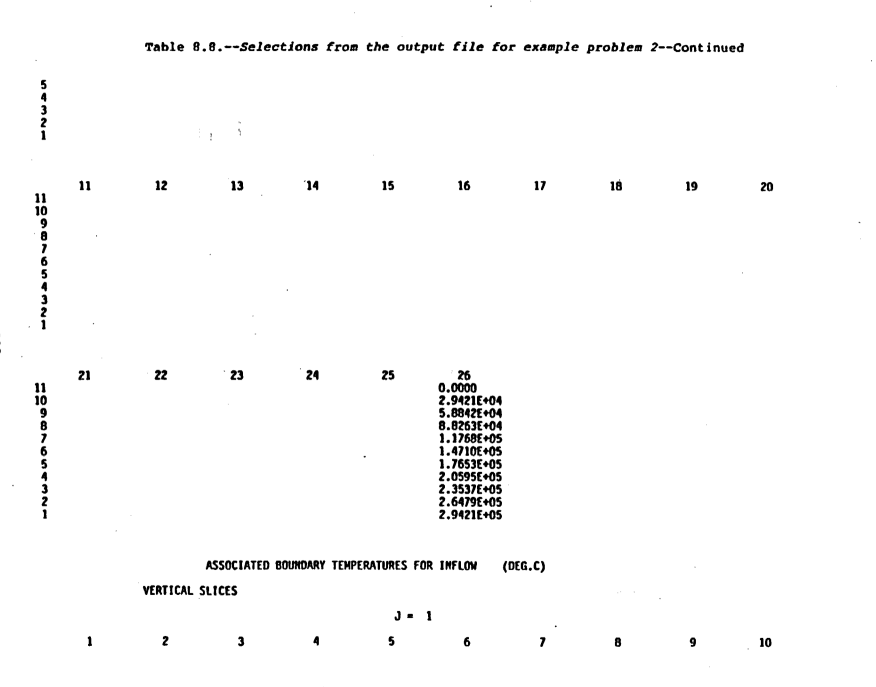
INDEX NUMBERS FOR SPECIFIED P,T OR C NODES

VERTICAL SLICES

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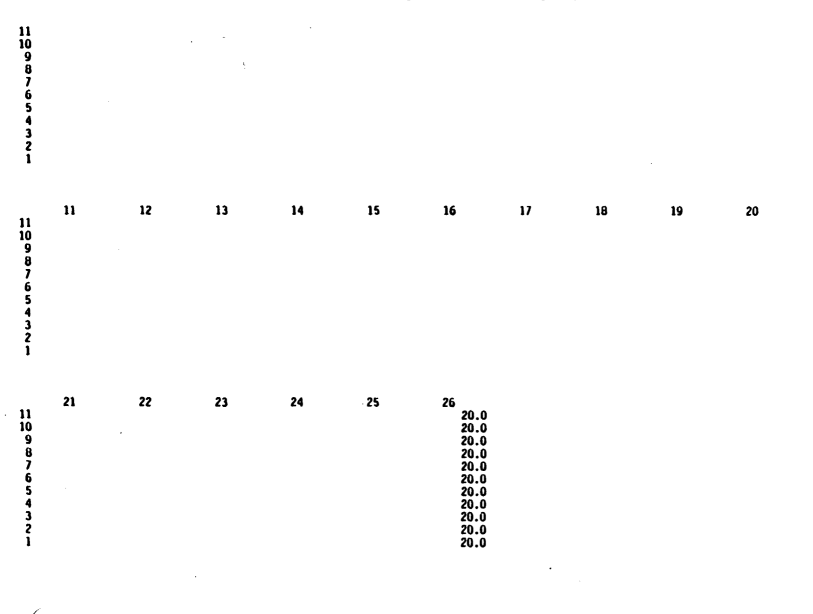


Table 8.8.--Selections from the output file for example problem 2--Continued

Table 8.8.--Selections from the output file for example problem 2--Continued \*\*\* TRANSIENT WELL DATA \*\*\* SURFACE WELL DATUM PRESSURE INJECTION INJECTION WELL FLOW RATE (M\*\*3/S) PRESSURE PRESSURE LIMITED? TEMPERATURE MASS FRACTION NO. ( PA) (PA) (DEG.C) (-) 60.00 1 0.2030 NO \*\*\* CALCULATION INFORMATION \*\*\* 3.00 (D) TIME AT WHICH NEXT SET OF TIME VARYING 90.0 (D) \*\*\* MAPPING DATA \*\*\* LENGTH OF Y-AXIS ..... 0.0 (IN.) LENGTH OF Z-AXIS ..... 6.0 (IN.) Z-AXIS IS POSITIVE UPWARD MAP NO. 11 12 J1 J2 K1 K2 MINIMUM VALUE MAXIMUM VALUE NUMBER OF OF VARIABLE OF VARIABLE ZONES 20. 60. A 2 1 26 1 1 1 11 +++ OUTPUT AT END OF TIME STEP NO. 30 \*\*\* 90.0 (D) 3.00 (D) NO. OF P.T.C LOOP ITERATIONS USED ..... 2 TEMPERATURE (DEG.C)

VERTICAL SLICES

J = 1

11 10 9 8 7 6 5 4 3 2 1	$ \begin{array}{c} 1 \\ 60.00 \\ 60$	2 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	3 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	4 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	5 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	6 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	7 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	8 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	9 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	$     \begin{array}{r}       10 \\       60.00 \\       6$
11 10 9 8 7 6 5 4 3 2 1	$ \begin{array}{c} 11 \\ 60.00 \\ 6$	12 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	13 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	14 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	$\begin{array}{c} 15 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \end{array}$	$\begin{array}{c} 16 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \\ 60.00 \end{array}$	17 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00 60.00	18     60.00     60	19 60.02 60.03 60.04 60.05 60.05 60.05 60.05 60.05 60.04 60.03 60.02	20 59.98 59.96 59.84 59.75 59.63 59.63 59.62 59.65 59.65 59.69 59.71
11 10 9 8 7 6 5 4 3 2 1	21 59.55 59.61 59.80 60.12 60.55 61.01 61.42 61.69 61.77 61.67 61.49	22 62.16 62.21 62.28 62.19 61.78 60.94 59.64 57.90 55.79 53.53 51.54	23 62.44 61.63 59.93 57.52 54.52 51.07 47.30 43.42 39.70 36.57 34.69	24 43.03 41.20 38.90 36.34 33.70 31.16 28.88 27.01 25.69 25.11	25 25.49 25.17 24.66 24.05 23.41 22.80 22.25 21.79 21.43 21.19 21.10	26 21.09 21.02 20.91 20.79 20.66 20.54 20.44 20.35 20.29 20.24 20.23				

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*** GLOBAL FLOW BALANCE SUMMARY *** CURRENT TIME STEP	RATES		AMOUNTS
FLUID INFLOW FLUID OUTFLOW CHANGE IN FLUID IN REGION RESIDUAL IMBALANCE FRACTIONAL IMBALANCE	2.014491E+02 -1.493657E+00 4.905415E-04	(KG/S) (KG/S) (KG/S) (KG/S)	5.182833E+07 (KG) 5.221562E+07 (KG) -3.871558E+05 (KG) 1.271484E+02 (KG) 0.0000
HEAT INFLOW HEAT OUTFLOW CHANGE IN HEAT IN REGION RESIDUAL IMBALANCE FRACTIONAL IMBALANCE	1.732711E+07 3.278493E+07 5.465375E+02	( J/S) ( J/S) ( J/S) ( J/S)	1.298890E+13 ( J) 4.491186E+12 ( J) 8.497854E+12 ( J) 1.416625E+08 ( J) 0.0000

## CUMULATIVE SUMMARY

AMOUNTS

FLUID INFLOW	1.554850E+09	(KG)
FLUID OUTFLOW	1.566620E+09	
CHANGE IN FLUID IN REGION	-1.176905E+07	(KG)
FLUID IN REGION	1.984422E+09	(KG)
RESIDUAL IMBALANCE	8.228376E+02	(KG)
FRACTIONAL IMBALANCE	0.0000	• •

HEAT INFLOW HEAT OUTFLON CHANGE IN HEAT IN REGION HEAT IN REGION RESIDUAL IMBALANCE	3.8966692+14	( )
CHANGE IN HEAT IN REGION	2.584199E+14	$\begin{pmatrix} 1\\ 1 \end{pmatrix}$
HEAT IN REGION	5.898154E+14 3.271183E+09	$\begin{pmatrix} 1\\ 1 \end{pmatrix}$
FRACTIONAL IMBALANCE	0.0000	

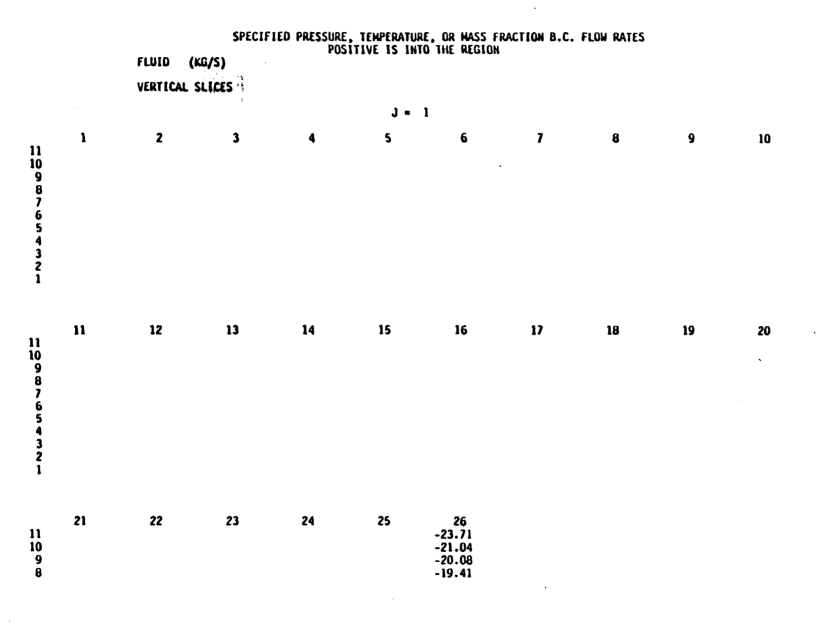
CUMINLATIVE SPECIFIED P CELL FLUID NET INFLOW	-1.566620E+09	(KG)
CUMULATIVE FLUX B.C. FLUID NET INFLOW	0.000000E-01	(KG)
CUMULATIVE LEAKAGE B.C. FLUID NET INFLOW	0.000000E-01	(KG)
CUMULATIVE AQUIFER INFLUENCE FLUID NET INFLOW	0.000000E-01	(KG)

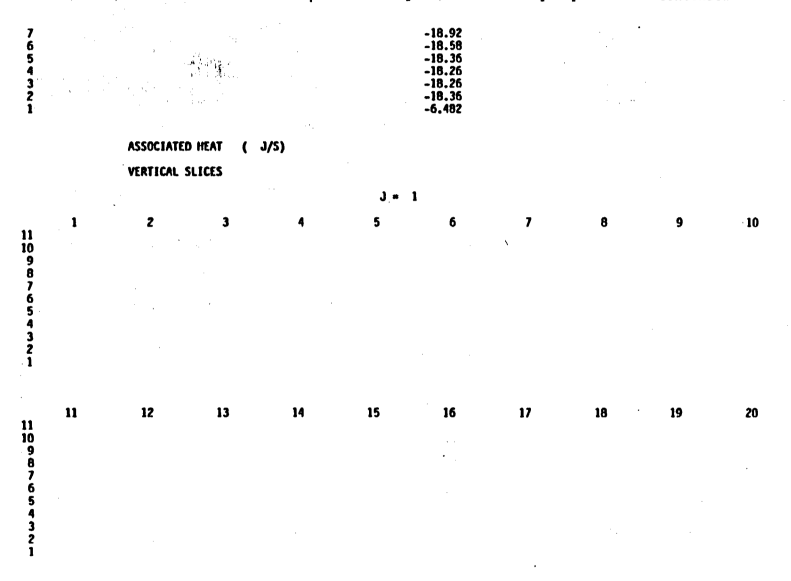
# CUMULATIVE SPECIFIED T CELL OR ASSOCIATED WITH

SPECIFIED P CELL HEAT NET INFLOW	-1.312503[+14	(J)
CUMULATIVE FLUX B.C. HEAT NET INFLOW	0.000000E-01	<i>i i i</i>
CUMULATIVE LEAKAGE B.C. HEAT NET INFLOW		
CUMULATIVE AQUIFER INFLUENCE HEAT NET INFLOW	0.000000E-01	2 35
CUMULATIVE HEAT CONDUCTION B.C. HEAT NET INFLOW	0.000000E-01	Ì

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	21	22	23	24	25	26
11						-2.0832E+06
10						-1.8430E+06
9			¢.			-1.7498E+06
8						-1.6810E+06
7						-1.6288E+06
6						-1.5904E+06
5						-1.5640E+06
4						-1.5484E+06
3						-1.5434E+06
2						-1.5489E+06
1						-5.4633E+05

VERTICAL SLICES

HEAT (J/S)

VERTICAL SLICES

VERTICAL PLANE AT ROW NO. 1

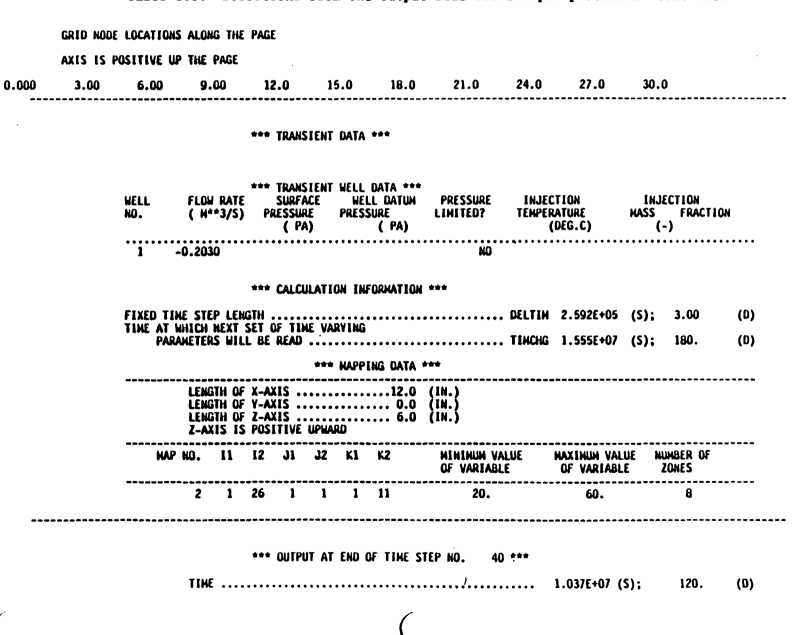
VERTICAL CROSS-SECTION OF SIMULATION REGION -	TENPERATU	RES	•	PAGE 1
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*44444444444444444444 <b>44444</b> 44444444444	33333	22222 1111	111
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	3333 22	22 111111	111
	333 222		
	333 2222		
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	222222	···· <b>111111111111</b>	
*4444444444444444444444444444444444444++++	222222	111111111111	
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	222222	111111111111	
	2222	111111111	
		1111111111	
444444444444444444444444444444444444444		111111111	
444444444444444444444444444444444444444		11111111	
*44444444444444444444444444444444444+++++	2 111	111111	
44444 <b>4444444</b> 444444444444444444444444++++++++	111	111111	
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****4*4*4*4*444*444*4444*4444*4++++*+44444 * 333 222*	1111		

NAP LEGEND HORIZONTAL GRID NODE RANGE, FROM 1 TO 26 VERTICAL GRID NODE RANGE, FROM 1 TO 11 DEPENDENT VARIABLE RANGE MAP CHARACTER

1
2
3
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	GRID NODE	LOCATIONS	ACROSS TH	E PAGE							
	1.25					3.75	4.67	5.82	7.26	9.04	11.3
14.0	17.5	21.8	27.2	33.9	42.3	52.7	65.6	81,8	102.	127.	158.
197.	246.										



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CURRENT TIME STEP LENGTH	2.592E+05 (S);	3.00	(D)
NO. OF P.T.C LOOP ITERATIONS USED	-3.3222E+02 ( PA)	AT LOCAT	ION ( 1, 1,11)
MAXIMUM CHANGE IN PRESSURE		) AT LOCAT	ION (23, 1,11)

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#### TEMPERATURE (DEG.C)

VERTICAL SLICES

VERTICAL SLICES							gar da g			
	· · ·				J = 1		:			
	1	2	3	4	5	6	7	8	9	10
11	59.22	59.22	59.22	59.22	59,22	59.22	59.22	59.23	59.24	59.24
10	59.17	59.17	59.17	59.17	59.17	59.17	59.17 59.02	59.18	59.18	59.19
9	59.02	59.02	59.02	59.02	59.02	59.02	59.02	59.02	59.03	59.04
9 . 8	58.77	58.77	58.77	58.77	58.78	58.78	58.78	58.78	58.78	58.79
7.	58.46	58.46	58.46	58.46	58.46	58.46	58.46	58.46	58.46	58.46
6	58.09	58,09	58.09	58.09	58.09	58.09	58.09	58.09	58.09	58.08
5	57.70	57.70	57.70	57.70	57.70	57.70	57.70	57.70	57.69	57.68
	57.33	57.33	57.33	57.33	57.33	57.70 57.33	57.32	57.32	57.31	57.29
3	57.02	57.02	57.02	57.02	57.02	57.02	57.01	57.00	56.99	56.97
4 3 2	56.81	56.81	56.81	56.81	56.81	56.80	56.80	56.79	56.77	56.74
1	56.73	56.73	56.73	56.73	56.73	56.73	56.72	56.71	56.69	56.66
		a server	· · · ·							
	11	12	13	14	15	16	17	18	19	20
11	59.25	59.27	59.28	59.29	59.30	59.29	59.26	59.18	59.01	58.62
10	59.20	59.21	59.23	59.23	59.24	59.22	59.18	59.09	58.88	58.43
9 8 7	59.04	59.05	59.06	59.06	59.05	59.02	58.95	58.81	58.52	57.91
8	58.79	58.79	58.79	58.78	58.75	58.69 58.26 57.74 57.19	58.58	58.36	57.94	57.09
7.	58.46	58.45	58.44	58.41	58.36	58.26	58.08	57.76	57.17	56.02
6	58.07	58.06	58.03	57.98	57.89	57.74	57.49	57.05	56.25	54.75
5	57.66	57.63	57.59	57.51	57.39	57.19	56.85	56.27	55.24	53.36
6 5 4 3	57.27	57.23	57.16	57.06	56.90	55.63	56.20	55.48	54.22	51.95
3	56.93	56.88	56.80	56.67	56.47	56.15	55.63	54.77	53.29	50.66
2	56.70	56.64	56.54	56.40	56.17	55.80	55.21	54.24	52.56	49.65
1	56.62	56.55	56.45	56.29	56.04	55.65	55.02	53.97	52.18	49.09
								. •		
••	21	22	23	24	25	26				
11	57.71	55.38	48.87	33.85	22.08	20.07	•			
10	57.38	54.67	47.43	32.84	21.94	20.00				

9	56.54	53.20	45.16	31.41	21.70	19.98
8	55.27	51.14	42.46	29.83	21.45	19.97
7	53.65	48.70	39.62	28.27	21.20	19.98
6	51.79	46.05	36.82	26.82	20.99	19.98
5	49.79	43.37	34.22	25.55	20.81	19.98
4	47.82	40.88	31.96	24.50	20.67	19.98
3	46.06	38.78	30.18	23.71	20.58	19.98
2	44.71	37.29	29.03	23.23	20.51	19.98
1	43.97	36.58	28.57	23.05	20.49	19.98

PAGE 1

VERTICAL PLANE AT ROW NO. 1

VERTICAL	CROSS-SECTION	OF SIMULATION REGION	- TEMPERATURES

33\*3333 22222 \* ÷ 22277.22 

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*44444444444444444444444444444444444444	333333333	222222	111111111
444444444444444444	333333333	2222222	11111111111
4444444444444444444444	333333333	2222222	111111111111
*4444444444444444	33333333	2222222	111111111111
4444444444444444	3333333	2222222	1111111111111
4444444444444444	3333333	2222222	1111111111111
4444444444444444	333333333	2222222	111111111111
*444444444444444	3333333	2222222	11111111111111
44444444444444	33333333	2222222	11111111111111
4444444444444444	3333333	2222222	11111111111111
****4**4*4*444* * *	3*333333 *	2222*22	11*111111111

MAP LEGEND HORIZONTAL GRID NODE RANGE, FROM 1 TO 26 VERTICAL GRID NODE RANGE, FROM 1 TO 11 DEPENDENT VARIABLE RANGE MAP CHARACTER

2.0008+01 - 2.50084	+01
2.500E+01 - 3.000E	01 1
3.000E+01 - 3.500E4	+01
3.500E+01 - 4.000E	H01 2
4.000E+01 - 4.500E	+01
4.500E+01 - 5.000E	+01 3
5.000E+01 - 5.500E	+01
5.500E+01 - 6.000E	H01 4

	GRID NOD	E LOCATIONS	i across th	e page							
1.00	1.25	1.55	1.94	2.41	3.01	3.75	4.67	5.82	7.26	9.04	11.3
14.0	17.5	21.8	27.2	33.9	42.3	52.7	65.6	81.8	102.	127.	158.
197.	246.										

GRID NODE LOCATIONS ALONG THE PAGE

AXIS IS POSITIVE UP THE PAGE

0.000 3.00 6.00 9.00 12.0 15.0 18.0 21.0 24.0 27.0 30.0

## \*\*\* OUTPUT AT END OF TIME STEP NO. 60 \*\*\*

TIME	1.555E+07 (S);	180.	(D)
CURRENT TIME STEP LENGTH	2.592E+05 (S);	3.00	(D)

NO. OF P,T,C LOOP ITERATIONS USED ..... 2 

TEMPERATURE (DEG.C)

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VERTICAL SLICES

J = 1

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	1	2	3	4	5	6	7	8	9	10
11	39.59	39.59	39.59	39.59	39.59	39.59	39.60	39.61	39.62	39.63
10	39.48	39.48	39.48	39.48	39.49	39.49	39.49	39.50	39.51	39.52
9	39.18	39.18	39.18	39.18	39.18	39.18	39.19	39.19	39.20	39.21
9 8 7	38.70	38.70	38.70	38.71	38.71	38.71	38.71	38.71	38.72	38.72
7	38.11	38.11	38.11	38.11	38.11	38.11	38.11	38.11	38.11	38.10
6	37.45	37.45	37.45	37.45	37.45	37.45	37.44	37.44	37.43	37.42
6 5	36.79	36.79	36.79	36.79	36.78	36.78	36.78	36.77	36.76	36.74
4	36.19	36.19	36.19	36.19	36.19	36.18	36.18	36.17	36.15	36.12
3	35.72	35.72	35.72	36.19 35.71	35.71	35.71	35.70	35.69	35.67	35.64
3 2	35.41	35.41	35.41	35.41	35.41	35.40	35.39	35.38	35.36	35.32
ī	35.31	35.31	35.31	35.30	35.30	35.30	35.29	35.27	35.25	35.21
-					•••••				30120	55.21
	11	12	13	14	15	16	17	18	19	20
11	39.65	39.66	39.66	39.64	39.59	39.46	39.21	38.76	37.99	36.72
10	39.54	39.55	39.55	39.52	39.46	39.32 38.93	39.06	38.60	37.82	36.52
9 8 7	39.22	39.22	39.21	39.17	39.09	38.93	38.65	38.15	37.32	35.98
8	38.72	38.71	38.68	38.63	38.53	38.34	38.01	37.46	36.58	35.18
7	38.09	38.07	38.03	37.95	37.82	37.59	37.22	36.62	35.67	34.21
6 5 4	37.40	37.36	37.30	37.20	37.04	36.77	36.35	35.70	34.69	33.18
5	36.71	36.66	36.58	36.45	36.26	35.96	35.50	34.79	33.73	32.18
4	36.08	36.02	35.93	35.78	35.56	35.23	34.73	33.99	32.89	31.31
3	35.59	35.52	35.41	35.25	35.01	34.66	34.13	33.35	32.22	30.62
2	35.27	35.19	35.08	34.91	34.66	34.29	33.74	32.95	31.80	30.19
1	35.16	35.08	34.96	34.79	34.54	34.16	33.61	32.81	31.65	30.04
	- 1		00		<b>0</b> 5					
	21	22	23	24	25	26				
11	34.68	31.54	27.13	22.21	19.60	19.94				
10	34.46	31.32	26.96	22.14	19.60	19.93				
9	33.88	30,75	26.51	21.97	19.62	19.92				

8	33.04	29.94	25.89	21.75	19.65	19.93
7	32.04	29.00	25.20	21.51	19.68	19.95
6	30.99	28.04	24.52	21.27	19.72	19.96
5	30,00	27.15	23.90	21.06	19.76	19.96
- 4	29.14	26,39	23.39	20.89	19.80	19.97
3	28.47	25,81	23.00	20.77	19.82	19.97
2	28.05	25.45	22.76	<b>20.69</b>	19.84	19.97
1	27.90	25.32	22.67	20.67	19.85	19.97

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	*** GLOBAL FLOW BALANCE SUMMARY *** CURRENT TIME STEP	RATES		AMOUNTS	
FLUID OUTFLOW CHANGE IN FLUID IN RESIDUAL IMBALANCE FRACTIONAL IMBALAN	CE	. 2.015989E+02 . 6.675322E-01 5.618930E-05	(KG/S) (KG/S) (KG/S) (KG/S)	5.242748E+07 5.225444E+07 1.730243E+05 -1.456427E+01 0.0000	(KG) (KG) (KG) (KG)
HEAT OUTFLOW CHANGE IN HEAT IN RESIDUAL INBALANCE	REGION	. 3.154910E+07 1.469188E+07 . 1.570947E+03	( J/S) ( J/S) ( J/S)	4.368984E+12 8.177527E+12 -3.808136E+12 4.071895E+08 0.0000	( J) ( J) ( J)

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### CUMULATIVE SUMMARY ٠

## AMOUNTS

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FLUID INFLOW FLUID OUTFLOW CHANGE IN FLUID IN REGION FLUID IN REGION RESIDUAL IMBALANCE FRACTIONAL IMBALANCE	3.128713E+09 -2.415865E+06 1.993776E+09 3.698014E+03	(KG) (KG) (KG) (KG) (KG)
HEAT INFLOW HEAT OUTFLOW CHANGE IN HEAT IN REGION HEAT IN REGION RESIDUAL IMBALANCE FRACTIONAL IMBALANCE	4.673636E+14 5.326113E+13 3.846566E+14 3.332536E+09	( J) ( J) ( J) ( J)
CUMULATIVE SPECIFIED P CELL FLUID NET INFLOW		(KG) (KG)

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CUMULATIVE SPECIFIED T CELL OR ASSOCIATED WITH

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#### \*\*\* WELL SUMMARY \*\*\*

WELL I NO. I			FLUID	G (POSITIVE HEAT ( J/S)	SOLUTE	TION) CUMULAT FLUID (KG)	HEAT SOLU	TE FLUID	IVE INJECTION HEAT SOLUTE (J) (KG)
1 1	1	1-11					9 3.36E+14		) 3.90E+14
		DUCTION	202. 0.000	3.155E+0 0.000	7	1.56E+09	9 3.36E+14	1.55E+09	3.90E+14
WELL NO.	TOP	CELL P	THE FOLLOG TION LAYER PRESSURE PA)	NELL DAT PRESSU	TUM	IN EFFECT DURIN WELL HEAD PRESSURE ( PA)	WELL DATUM	WELL HEAD TEMPERATURE	NASS FRACTION (-)
					********			0.0	
1	PE	-4.187 Er layef		-4.1870		TES- (KG/S) (IH.	38.4 JECTION IS POSIT		
		ER LAYEF	FLUID PROL		ECTION RA	TES- (KG/S) (IN			
		ER LAYEF	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN.			
AYER 1 11 10	NO. - 10 - 21	1 0.5 1.0	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN.			
AYER #	NO. - 10 - 21 - 20	1 0.5 1.0 0.9	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN			
AYER   11 10 9 8 7	NO. - 10 - 21	1 0.5 1.0 0.9 0.7	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN			
LAYER   11 10 9 8 7	NO. - 10 -21 -20 -20 -20 -20	1 0.5 1.0 0.9 0.7 0.4 0.2	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN.			
LAYER ) 11 10 9 8 7 6 5	NO. - 10 - 21 - 20 - 20 - 20 - 20 - 19	1 0.5 1.0 0.9 0.7 0.4 0.2 9.9	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN			
AYER 1 11 10 9 8 7	NO. - 10 -21 -20 -20 -20 -20	1 0.5 1.0 0.9 0.7 0.4 0.2 9.9 9.6	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN			
LAYER   11 10 9 8 7 6 5 4	NO. - 10 - 21 - 20 - 20 - 20 - 20 - 19 - 19	1 0.5 1.0 0.9 0.7 0.4 0.2 9.9 9.6 9.4	FLUID PROL	DUCTION/INJ	ECTION RA	TES- (KG/S) (IN			

1 -9.64

	PER LAYER HEAT PRODUCTION/INJECTION RATES- ( J/S) (INJECTION IS POSITIVE)						
LAYER	NO. 1	WELL NUMBER					
11 10 9 8 7 6 5 4 3 2 1	-1.739E+06 -3.461E+06 -3.414E+06 -3.340E+06 -3.249E+06 -3.150E+06 -3.053E+06 -2.967E+06 -2.899E+06 -2.856E+06 -1.421E+06						

PAGE 1 VERTICAL CROSS-SECTION OF SIMULATION REGION -TEMPERATURES 1111111+111111 \_\_\_\_\_ \*22222222222222222 \_\_\_\_\_\* -----\* \*222222222222222222222222 

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## 9. NOTATION

9.1 ROMAN

- a coefficients of the various u ijk in the spatially discretized equations.
- a coefficients of the various uik in the spatially discretized equations for the cylindrical-coordinate system.
  - a1 implicit term for aquifer-influence-function or heat-conduction boundary conditions (appropriate units).
  - a2 implicit term coefficient for aquifer-influence-function or heat-conduction boundary conditions (appropriate units).
  - <u>A</u> matrix of coefficients for the <u> $\delta u$ </u> vector for the finite-difference equations (eq. 3.6.2.1a).
- $\underline{A}_i$  Banded, sparse submatrices of the matrix  $\underline{A}$  for the finitedifference equations.
- A capacitance coefficient (appropriate units).
- $A_{ms}$  value of capacitance coefficient within subdomain s of cell m.
  - b vector of known terms on the right-hand side for the finitedifference equations.
- b\_ thickness of an outer-aquifer region (m).
- b<sub>HC</sub> effective thickness of a heat-conducting medium exterior to the region (m).

b, thickness of an aquitard layer (m).

b<sub>p</sub> thickness of a riverbed sediment (m).

b<sub>i</sub> coefficient in the approximation to the transient aquifer-influence function given by equation 3.4.4.2.5.

B coefficient of dispersion or diffusion (appropriate units).

- $B_0, B_1$  parameters for the temperature-dependence of viscosity (°C).
  - B tensor of diffusion or dispersion of rank 3 (appropriate units).
  - $B_{ij}$  tensor components of  $\underline{B}$ .
    - $\overline{c_f}$  average heat capacity of the fluid phase at constant pressure  $(J/kg^{-o}C)$ .
  - $c_{s_{n}}$  heat capacity of pure water at constant pressure (J/kg-°C)
  - c heat capacity of the exterior heat-conducting medium at constant pressure (J/kg-°C).
  - C coefficient in the capacitance matrix for the discretized equations (appropriate units).
    - C vector of interstitial velocity (m/s).
    - $C_i$  (m,p,q) vector components of <u>C</u> in the ith direction, for the cell m, face p, in element q (m/s).
    - d input data item.
  - d coefficient in the series expansion for pressure (eq. 3.3.2.4c) (Pa/m).

- d<sub>Ti</sub> coefficient in the series expansion for temperature (eq. 3.3.2.4d) (°C/m).
  - d coefficient vector for series expansion (eq. 3.3.2.5b).
  - $\underline{D}_i$ , submatrices on the diagonal of matrix  $\underline{A}$ .

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- $D_m$  effective-molecular-diffusivity coefficient of the solute  $(m^2/s)$ .
- $D_{Hrm}$  thermal diffusivity of the medium surrounding a well riser (m<sup>2</sup>/s).
  - $\underline{\mathbf{D}}_{\mathbf{W}}$  thermo-mechanical dispersion tensor from flow mechanisms (W/m-°C).
- $\begin{array}{c} D_{\text{He}} \\ \text{boundary } (m^2/s). \end{array}$
- $D_{\text{Hij}}$  thermo-mechanical-dispersion-tensor component (W/m-°C).
  - $\underline{P}_{S}$  mechanical-dispersion-coefficient tensor  $(m^2/s)$ .
- $D_{Sii}$  mechanical-dispersion-tensor component (m<sup>2</sup>/s).
- $D_{Sij}^{*}$  hydrodynamic-dispersion coefficient (m<sup>2</sup>/s).
- $D_{\text{Hij}}^{\star}$  thermo-hydrodynamic-dispersion coefficient (W/m-°C).
  - D coefficient for a source proportional to the dependent variable (appropriate units).
- $D_{ms}$  value of D in cell m for subdomain s.

- $\underline{E}_i$  coefficient vector for a node; i = 1 for flow equation, i = 2 for heat-transport equation, i = 3 for solute-transport equation.
  - E source-term intensity (appropriate units).
- $E_{s(m)}$  source-term intensity for source s in cell m (appropriate units).
  - E value of source term in cell m.
  - fraction of cell thickness that is saturated for a free-surface boundary condition (-).
  - f\_ head-loss friction factor for a well riser (-).
  - f. head-loss friction factor.for a well bore (-).
  - f<sub>0</sub> angle-cf-influence factor for aquifer-influence-function boundary condition (-).
  - F<sub>CJ</sub> heat-flux function (Carslaw and Jaeger) to an infinite medium from a constant-temperature cylindrical source (-).
  - $F_i$  vector component of the known terms at time level n in the three discretized system equations for a given node, i = 1,2,3.
  - F spatial finite-difference function.
  - $F^{II}$  spatial finite-difference function evaluated at time level n.
  - $F_{CJ}^{S}$  approximation to  $F_{CJ}$  function for small dimensionless time.
  - $F_{C,I}^{L}$  approximation to  $F_{C,J}$  function for large dimensionless time.
  - F function for the well-riser calculation (eq. 2.4.2.11).

g gravitational constant  $(m/s^2)$ .

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g component of gravitational acceleration in the direction normal to the face p.

G function for the well-riser calculation (eq. 2.4.2.11).

h, heat-transfer coefficient from the fluid to a riser pipe  $(W/m^2-{}^{\circ}C)$ .

H specific enthalpy of the fluid phase (J/kg).

 $H_{p}$  specific enthalpy of fluid at a region boundary (J/kg).

 $H_{a}$  specific enthalpy of fluid in an outer aquifer (J/kg).

H\_ specific enthalpy of fluid in a well riser (J/kg).

H specific enthalpy of the solid phase (J/kg).

J vector of specified total flux at a boundary (appropriate units).

 $J_{i}$  component of specified flux at a boundary in the ith direction.

J component of specified flux at a boundary in the ith direction in element q.

k porous-medium permeability tensor (m<sup>2</sup>).

k permeability of an outer-aquifer region  $(m^2)$ .

 $k_{T}$  permeability of an aquitard layer (m<sup>2</sup>).

 $k_{R}$  permeability of a riverbed sediment (m<sup>2</sup>).

k, mean permeability around a well between  $r_{u}$  and  $r_{e}$  ( $m^{2}$ ).

k permeability in the x-direction  $(m^2)$ .

 $k_v$  permeability in the y-direction (m<sup>2</sup>).

 $k_{pq}$  permeability in the direction normal to face p for subdomain q (m<sup>2</sup>).

 $k_{wm(l)}$  average radial permeability around the well in cell m at level  $l(m^2)$ .

 $k_{mpq}$  permeability for cell m, face p, subdomain q (m<sup>2</sup>).

 $K_{A}$  equilibrium-distribution coefficient (m<sup>3</sup>/kg).

K thermal conductivity of a medium exterior to the simulation region  $(W/m^{\circ}C)$ .

 $K_{f}$  thermal conductivity of the fluid (W/m-°C).

 $K_r$  thermal conductivity of a riser pipe (W/m-°C).

K<sub>r</sub> thermal conductivity of the medium surrounding a riser pipe ( $W/m^{-o}C$ ).

K thermal conductivity of the solid phase (porous matrix) ( $W/m-^{\circ}C$ ).

K Augmented porosity factor for subdomain s (eq. 3.1.4.4k) (-).

l distance along the well bore or well riser (m).

 $l_{T}$  lower end of a well-screen interval (m).

 $\hat{\boldsymbol{z}}_{T}$  index of the bottom level of a well screen.

 $l_{11}$  upper end of a well-screen interval (m).

 $\hat{l}_{11}$  index of the top level of a well screen.

		L_	length of a well riser (m).						
$\bigcirc$	)	L <sub>L1</sub>	length of a well bore in the lower half of cell m at level $\ell$ (m).						
		L <sub>22</sub>	length of a well bore in the upper half of cell m at level $\ell$ (m).						
		<u>L</u> i	submatrices below the diagonal of matrix A.						
		n	cell number with coordinate indices i,j,k.						
		m(i-1)	cell number with coordinate indices i-1,j,k.						
		m(£)	cell number associated with well bore level £.						
		М	total number of nodes in the simulation region.						
		M <sub>A</sub>	number of pot-aquifer boundary-condition cells.						
$\bigcirc$		Mw	well mobility per unit length of well bore (m <sup>3</sup> /s-m-Pa).						
	•.	Mwl	well mobility per unit length of well bore at level $\ell$ (m <sup>3</sup> /s-m-Pa).						
		Mn	mass of fluid plus effective additional fluid mass from sorption in cell m (kg).						
		n	data repeat factor.						
		n	vector in the outward normal direction to the boundary.						
		N <sub>r</sub>	number of grid points in the r-direction.						
		N <sub>x</sub>	number of grid points in the x-direction.						
		N y	number of grid points in the y-direction.						

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- $N_{\perp}$  number of grid points in the z-direction.
- N number of line sources in the region.
- p fluid pressure relative to atmospheric pressure (Pa).
- p̂ absolute pressure (Pa).
- $p_{av}$  average pressure around a well between  $r_{av}$  and  $r_{av}$  (Pa).
- p<sub>R</sub> pressure at a boundary (Pa).
- p dimensionless pressure at an aquifer-influence-function boundary (-)
   (eq. 2.5.4.2.6b).

p\_ pressure in the outer-aquifer region (Pa).

 $p_{a}^{0}$  initial specified pressure in an outer-aquifer region (Pa).

p<sub>inj</sub> pressure of injected fluid (Pa).

- p<sub>m</sub> pressure at node m (Pa).
- p<sub>o</sub> pressure at a reference state for density (Pa).

 $p_{oH}$  pressure at a reference state for enthalpy (Pa).

- $\hat{p}_{oH}$  absolute pressure at a reference state for enthalpy (Pa).
- p, pressure averaged across a riser cross section (Pa).
- $p_{rk}$  pressure averaged across a riser cross section at location  $\ell_k$  (Pa).

- $\hat{p}_{sat}$  absolute pressure of saturated water at zero degrees Celsius (Pa).  $p_{sat}$  pressure at the well bore (Pa).
- P<sub>wd</sub> pressure at a well datum (Pa).
- $P_{w\ell}$  pressure in the well bore at level  $\ell$  (Pa).
- $p^0$  initial pressure distribution (Pa).
- P U dimensionless pressure response to a unit-step withdrawal flow rate (-).
  - q fluid-source, flow-rate intensity  $(m^3/m^3-s)$ .
- $q_{F_i}^B$  specified fluid-flux-vector components  $(m^3/m^2-s)$ .
- $q_{r_n}$  normal component of fluid flux  $(m^3/m^2-s)$ .
- $q_{fw}$  fluid flux from a well  $(m^3/m^2-s)$ .

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- q<sub>H</sub> heat-source-rate intensity (W/m<sup>3</sup>).
- $q_{HC}$  heat flux at the heat-conduction boundary (W/m<sup>2</sup>).

 $q_{Hi}^{B}$  specified heat-flux-vector components (W/m<sup>2</sup>).

 $q_{\rm HL}$  heat flux across a leakage boundary (W/m<sup>2</sup>).

 $q_{\mu_m}$  normal component of heat flux (W/m<sup>2</sup>).

- $q_r$  fluid flux across a leakage boundary  $(m^3/m^2-s)$ .
- $q_p$  fluid flux across a river-leakage boundary  $(m^3/m^2-s)$ .
- $q_{Rmax}$  limit on the fluid flux from a river to the aquifer  $(m^3/m^2-s)$ .
  - $q_{S_i}^B$  specified solute-flux-vector components (kg/m<sup>2</sup>-s).
  - $q_{ct}$  solute flux across a leakage boundary (kg/m<sup>2</sup>-s).
  - $q_{Sn}$  normal component of solute flux (kg/m<sup>2</sup>-s).
  - q<sub>w</sub> volumetric flow rate per unit length of well bore; (positive is from the well to the aquifer) (m<sup>3</sup>/m-s).
  - Q fluid-mass-source flow rate (kg/s).
  - Q'A dimensionless flow rate per unit thickness at an aquifer-influencefunction boundary (-).
  - Q<sub>A</sub> constant specified flow rate at an aquifer-influence-function boundary (m<sup>3</sup>/s).
  - Q<sub>Am</sub> volumetric flow rate across the boundary for cell m between the innerand outer-aquifer regions; (positive is into the inner region) (m<sup>3</sup>/s).
  - $Q_B$  specified flow rate per unit thickness at the aquifer-influencefunction boundary of the region (m<sup>3</sup>/m-s).
  - $Q_{Fr}$  mass flow rate in a well riser (kg/s).
- $Q_{\rm HCm}$  heat-flow rate across a heat-conduction boundary at cell m (W).

Q<sub>Hr</sub> heat transfered per unit mass per unit length to the fluid in a riser (J/kg-m).

- $Q_{rm}$  volumetric-flow rate across a leakage boundary at cell m (m<sup>3</sup>/s).
  - Q<sup>n</sup><sub>m</sub> volumetric-flow rate of a source term for cell m at time level n including specified-flux boundary condition and line sources (m<sup>3</sup>/s).
- $Q'_{11}$  dimensionless flow-rate response to a unit step change of pressure.
- $Q_{\rm Rm}$  volumetric-flow rate across a river-leakage boundary at cell m  $(m^3/s)$ .

Q. volumetric-flow rate from a well to the aquifer  $(m^3/s)$ .

 $Q_{w\ell}$  volumetric-flow rate from a well to the aquifer at well bore level  $\ell$  (m<sup>3</sup>/s).

r distance along the r-coordinate direction (m).

r radius of influence of a well (m).

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 $r_r$  exterior radius of an outer-aquifer region (m).

r<sub>1</sub> interior radius of the region, cylindrical coordinates (m).

 $r_{i+\frac{1}{2}}$  radius of the cell boundary between the node at  $r_i$  and the node at  $r_{i+1}$  (m).

r, interior radius of outer-aquifer region (m).

r<sub>N</sub> exterior radius of the region, cylindrical coordinates (m). r r\_ inner radius of a well-riser pipe (m).

- r, well-bore radius (m).
- R ratio of exterior to interior radius for an outer-aquifer region for the aquifer-influence-function boundary condition (-).
- R<sub>fs</sub> transfer of solute from fluid to solid phase per unit mass of solid phase (kg/s-kg).
- R right-hand side terms for the ith equation, i = 1,2,3 for pressure, temperature, and mass fraction respectively.
- $\frac{R}{m}$  rational-function vector that approximates the right-hand-side of equation 3.3.2.6.
- $R(L_1)$  spectral radius of the Gauss-Seidel iteration matrix.
  - S specific storage for a confined aquifer (eq. 2.2.5.5)  $(m^{-1})$ .
- S<sub>Alpq</sub> area of the aquifer-influence-function boundary face for cell l, face p, element q ( $m^2$ ).
  - S boundary surface of cell m  $(m^2)$ .
  - S part of the boundary surface of cell m that belongs to face  $p(m^2)$ .
  - S part of the boundary surface of cell m that belongs to face p in element  $q(m^2)$ .
- S part of the boundary surface of cell m that is an aquifer-influence-function boundary  $(m^2)$ .
- S part of the boundary surface of cell m that is a leakage boundary  $(m^2)$ .

 $S_{BHCm}$  part of the boundary surface of cell m that is a heat-conduction boundary  $(m^2)$ .

- S<sup>1</sup><sub>u</sub> parts of the region boundary where specified-value boundary conditions are applied (Dirchlet boundary conditions); u = p, T, or w.
- S<sup>2</sup><sub>u</sub> parts of the region boundary where specified-flux boundary conditions are applied (Neumann boundary conditions); u = p, T, or w.
- $S^3$  part of the boundary that is an aquifer-leakage boundary.

 $S^4$  part of the boundary that is a river-leakage boundary.

 $S^5$  part of the region boundary that is a heat-conduction boundary.

 $S^6$  part of the region boundary that is a free-surface boundary.

t time (s).

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t<sup>n</sup> time at level n (s).

t' dimensionless time defined by equation 2.5.4.2.5c (-).

T temperature of the fluid and porous medium (°C).

T\_ ambient temperature of the medium adjacent to a well riser (°C).

T<sub>p</sub> temperature at a boundary (°C).

Te temperature in the medium exterior to a heat-conduction boundary (°C).

T temperature at a reference state for density (°C).

- T<sub>oH</sub> temperature at a reference state for enthalpy (°C).
- T<sub>ov</sub> temperature at a reference state for fluid viscosity (°C).
- $T_{r}$  temperature of the fluid averaged across a riser cross section (°C).
- T<sub>1</sub> temperature solution to the first heat-conduction problem (eq. 2.5.5.3a-d) (°C).
- T<sub>2</sub> temperature solution to the second heat-conduction problem (eq. 2.5.5.4a-d) (°C).
- T\* temperature of a fluid source (°C).
- T<sup>0</sup> initial-temperature distribution (°C).
- $T_{\perp}^{0}$  initial-temperature profile in the exterior medium (°C).
- T<sub>ini</sub> temperature of injected fluid (°C).
  - T<sub>U</sub> unit-step temperature-response solution to heat-conduction problem (eq. 2.5.5.6b) (°C).
  - $T_{ud}$  temperature of the fluid at a well datum (°C).
    - **T** absolute temperature (K).
  - $T_{ri}$  conductance for flow (kg/s-Pa) or (m-s).
  - $T_{\mu_i}$  conductance for heat transport (W/°C).
  - $T_{si}$  conductance for solute transport (kg/s).
  - $T_{wF}$  conductance for flow at a well bore in the cylindrical coordinate system (m-s).

u generic dependent variable (appropriate units).

- $u_{\rm p}$  boundary-condition distribution of u.
- u<sup>0</sup> initial-condition distribution of u.
- $u_{ijk}^n$  value of u at  $x_i$ ,  $y_j$ ,  $z_k$  at time  $t^n$ .
- $u_{ik}^{n}$  value of u at  $x_{i}$ ,  $z_{k}$  in cylindrical-coordinate system at time  $t^{n}$ .
- $U_T$  overall heat-transfer coefficient for the fluid, riser-pipe, and surrounding medium ( $W/m^2-^{\circ}C$ ).

 $\underline{V}_i$  submatrices of matrix  $\underline{A}$  above the diagonal.

v interstitial-velocity vector (m/s).

v magnitude of the interstitial velocity (m/s).

 $v_i$  interstitial-velocity component in the ith direction (m/s).

v velocity averaged across a riser cross section at a given z-level
 (m/s).

v velocity averaged across a well-bore cross section at a given z-level
 (m/s).

V region of the aquifer for simulation.

 $V_{\rm b}$  bulk or total volume of a fixed mass of porous medium (m<sup>3</sup>).

 $V_{a}$  volume of an outer-aquifer region (m<sup>3</sup>).

- $V_{\underline{\ell}}$  volume of an outer-aquifer region that influences boundary cell  $\underline{\ell}$  (m<sup>3</sup>).
- $V_{m}$  volume of cell m (m<sup>3</sup>).
- $V_{me}$  volume of subdomain s in cell m (m<sup>3</sup>).
  - w mass fraction of solute in the fluid phase (-).
  - $w_n$  mass fraction of solute at a boundary (-).
  - we mass fraction of solute in an outer aquifer outside a leakage boundary (-).

w<sub>min</sub> minimum solute-mass fraction for scaling (-).

w maximum solute-mass fraction for scaling (-).

- $w_{n}$  mass fraction of solute at a reference state for density (-).
- w' scaled-mass fraction of solute defined by equation 2.2.1.2 (-).

w<sup>U</sup> mass fraction of solute initial distribution (-).

w\* mass fraction of solute in the fluid source (-).

 $\overline{w}$  mass fraction of solute on the solid phase (-).

- W cumulative net flow from an outer- to an inner-aquifer region  $(m^3)$ .
- $W_{I}$  well index per unit length of well bore defined by equation 2.4.1.2 (m<sup>2</sup>).

 $\underline{x}_{m}$  position vector of node point m (m).

- x distance along the x-coordinate direction (m).
- <u>x</u> vector of position in the simulation region including the boundary (m).

 $\underline{x}_{s(m)}$  location vector of line source s in cell m (m).

y distance along the y-coordinate direction (m).

Y Pressure and temperature vector in a well riser.

z distance along the z-coordinate or vertical direction (m).

z<sub>p</sub> elevation of the region boundary (m).

z\_ elevation of the top of an aquitard layer (m).

 $z_o$  elevation of the node in a well at level  $\ell$  (m).

z<sub>LS</sub> elevation of the land surface (m).

 $z_m$  elevation of the node of cell m (m).

 $z_{p}$  coordinate in the outward-normal direction to the boundary (m).

z<sub>p</sub> elevation of a river bottom (m).

z<sub>RFS</sub> elevation of the water surface of a river (m).

z<sub>wd</sub> elevation of a well datum (m).

9.2 GREEK

- $\alpha_{\rm b}$  bulk compressibility of the porous medium (Pa<sup>-1</sup>).
- $\alpha_{\rm be}$  bulk compressibility of the porous medium in an outer region (Pa<sup>-1</sup>).
- $\alpha_r$  dispersivity longitudinal to the flow direction (m).

 $\alpha_{\rm T}$  dispersivity transverse to the flow direction (m).

α rock compressibility (Pa<sup>-1</sup>).

- $\gamma_{Am}$  apportionment factor for aquifer-influence-function flow rate (-).
- $\gamma_p$  fraction of riverbed area per unit area of aquifer boundary (-).
- $\beta_{n}$  fluid compressibility (Pa<sup>-1</sup>).

 $\beta_{T}$  fluid coefficient of thermal expansion (°C<sup>-1</sup>).

 $\beta_{W}$  slope of fluid density as a function of mass fraction divided by the reference fluid density (-).

 $\beta_{\rm pe}$  fluid compressibility in an outer region (Pa<sup>-1</sup>).

 $\beta_{W}$  slope of fluid density as a function of scaled-mass fraction divided by the reference fluid density (-).

ε effective porosity (-).

 $\varepsilon_{a}$  effective porosity of an outer-aquifer region (-).

 $\lambda$  linear decay constant (s<sup>-1</sup>).

- $\lambda$  maximum estimate of spectral radius or maximum eigenvalue (eq. 3.7.2.6b).

 $\mu$  viscosity of the fluid (kg/m-s).

 $\mu_e$  viscosity of the fluid in an outer-aquifer region (kg/m-s).

 $\mu_{\tau}$  viscosity of fluid in an aquitard layer (kg/m-s).

 $\mu(T_{out}, w')$  viscosity at the reference temperature (kg/m-s).

 $\mu_p$  viscosity of the fluid in a riverbed (kg/m-s).

 $\bar{\mu}_{p}^{n}$  average viscosity at cell face p at time t<sup>n</sup> (kg/m-s).

 $\mu_0$  viscosity at the minimum mass fraction (kg/m-s).

 $\mu_1$  viscosity at the maximum mass fraction (kg/m-s).

 $\mu_r$  viscosity of the fluid in a riser pipe (kg/m-s).

v counter index for successive-overrelaxation iteration.

poténtial energy per unit mass of fluid in the aquifer outside
 a leakage boundary (Nt-m/kg).

 $\rho$  fluid density (kg/m<sup>3</sup>).

ρ<sub>b</sub> bulk density of the porous medium (mass/unit volume of dry porous medium) (kg/m<sup>3</sup>).

- $\rho_{\rm R}$  fluid density at the region boundary (kg/m<sup>3</sup>).
- $\rho_{\perp}$  fluid density outside a leakage boundary (kg/m<sup>3</sup>).
- $\rho_0$  fluid density at a reference pressure, temperature, and mass fraction (kg/m<sup>3</sup>).
- $\rho_{\_}$  fluid density averaged across a well-riser cross section (kg/m<sup>3</sup>).
- $\rho_R$  fluid density of the river (kg/m³).
- $\rho_s$  density of the solid phase of the porous matrix (mass/unit volume of solid phase) (kg/m<sup>3</sup>).
- $\rho_{se}$  density of the solid phase of the porous matrix exterior to the simulation region (mass/unit volume of solid phase) (kg/m<sup>3</sup>).
- $\rho_{\rm c}$  average fluid density in a well bore (kg/m<sup>3</sup>).
- $\bar{\rho}_{\rm p}^{\rm n}$  average density at cell face p at time t<sup>n</sup> (kg/m<sup>3</sup>).
  - $p^*$  density of a fluid source (kg/m<sup>3</sup>).
  - $\sigma$  spatial weighting coefficient (eq. 3.1.4.4).
  - t dimensionless time for heat transfer from a defined well riser by
    equation 2.4.2.6c (-).

 $\theta$  factor for time differencing (eq. 3.1.3.4).

- $\theta_{\perp}$  angle between the well riser and the z-coordinate (deg).
- $\theta_{ij}$  angle between the well bore and the z-coordinate (deg).

w overrelaxation factor (eq. 3.7.2.2).

w optimum overrelaxation factor (eq. 3.7.2.7).

9.3 MATHEMATICAL OPERATORS AND SPECIAL FUNCTIONS

 $\delta u$  vector of dependent variables for the finite-difference equations.

δu temporal change in u.

∆u spatial change in u.

δu maximum temporal change in u.

 $\delta u_{max}^{S}$  specified maximum temporal change in u.

ū average value of u.

2

 $\Delta r_r$  wall thickness of a well-riser pipe (m).

St current time-step length.

δt previous time-step length.

 $\nabla$  del operator;  $\hat{e}_x \frac{\partial}{\partial x} + \hat{e}_y \frac{\partial}{\partial y} + \hat{e}_z \frac{\partial}{\partial z}$  in cartesian coordinates;

 $\hat{e}_r \frac{\partial}{\partial r} + \hat{e}_{\theta} \frac{1}{r} \frac{\partial}{\partial \theta} + \hat{e}_z \frac{\partial}{\partial z}$  in cylindrical coordinates.

| | absolute value.

• • •

 $\frac{\partial}{\partial n}$  derivative in the outward-normal direction at a boundary.

 $\delta_{ij}$  Kronecker delta function; =1 for i=j, = 0 for i≠j.

 $\delta(\underline{x}-\underline{x}_{e})$  delta function for the point source in a cell at  $\underline{x}_{e}$ .

I identity matrix of rank 3.

J Bessel function of the first kind, order n.

Y Bessel function of the second kind, order n.

Y Euler's constant  $\simeq 0.577$ .

 $\partial p_b$  $\partial t$  $\partial t$  $\partial t$ influence-function boundary (Pa/s).

 $\frac{\partial \overline{p}_e}{\partial t}$  spatial average of the rate of pressure change in an outeraquifer region (Pa/s)

Dp bl

average rate of pressure change at the boundary of an inner region (Pa/s)

 $\frac{\partial Q_m}{\partial p} \Big|_n$  implicit term for fluid line sources at the nth time level (kg/s-Pa).

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## 11. SUPPLEMENTAL DATA

Because of the excessive length (about 12,000 lines), no program listing is provided for the HST3D code. However, listings of the major program variables and a cross-reference table are provided to aid the user in error tracing and program modification.

## 11.1 HST3D PROGRAM VARIABLE LIST WITH DEFINITIONS

The following program-variable list (table 11.1) contains all of the major variable names with brief definitions. Minor variables including loop indices, array subscripts, and temporary results, have not been included. Most temporary variables begin with the letter U with the remainder of the name based on the corresponding major variable. Variably partitioned array pointers begin with the letter I.

A1HC - HEAT CONDUCTION B.C. GEOMETRIC FACTOR A2HC - HEAT CONDUCTION B.C. GEOMETRIC FACTOR A3HC - HEAT CONDUCTION B.C., GEOMETRIC FACTOR A4 - SUB-MATRIX OF SYSTEM EQUATIONS FOR D4 SOLVER AAIF - AQUIFER INFLUENCE FUNCTION B.C. CONSTANT TERM ABOAR - POROUS MEDIUM COMPRESSIBILITY FOR OUTER AQUIFER REGION ABPM - POROUS MEDIUM COMPRESSIBILITY ALBC - LEAKAGE B.C. CONSTANT TERM ALPHL - LONGITUDINAL DISPERSIVITY ALPHT - TRANSVERSE DISPERSIVITY AMAX - UPPER LIMIT OF VARIABLE FOR CONTOUR MAP AMIN - LOWER LIMIT OF VARIABLE FOR CONTOUR MAP ANGOAR - ANGLE OF INFLUENCE FOR CARTER-TRACY AQUIFER INFLUENCE FUNCTION B.C. APLOT - OUTPUT ARRAY FOR CONIGUR MAP APRNT - OUTPUT LINE FOR ARRAY PRINTING ARGRID - TRUE IF AUTOMATIC RADIAL NODE LOCATIONS TO BE SET ARRAY - OUTPUT ARRAY FOR PRINTED TABLES ARWB - CROSS-SECTIONAL AREA OF WELL BORE ARX - CELL-FACE AREA IN Y-Z PLANE ARXBC - CELL-FACE AREA IN Y-Z PLANE FOR BOUNDARY CONDITIONS ARXFBC - CELL-FACE AREA IN Y-Z PLANE FOR FLUX B.C ARY - CELL-FACE AREA IN X-Z PLANE ARYBC - CELL-FACE AREA IN X-Z PLANE FOR BOUNDARY CONDITIONS ARYFBC - CELL-FACE AREA IN X-Z PLANE FOR FLUX B.C ARZ - CELL-FACE AREA IN X-Y PLANE ARZBC - CELL-FACE AREA IN X-Y PLANE FOR BOUNDARY CONDITIONS ARZFBC - CELL-FACE AREA IN X-Y PLANE FOR FLUX B.C AST - \*\*\* AUTOTS - TRUE IF AUTOMATIC TIME STEPPING IS DESIRED BAIF - AQUIFER INFLUENCE FUNCTION B.C. TEMPORAL COEFFICIENT BBLBC - THICKNESS OF CONFINING LAYER OR RIVER BED BLANK - ! ! BLANKL - 1 BLBC - LEAKAGE B.C. TEMPORAL COEFFICIENT BOAR - THICKNESS OF OUTER AQUIFER REGION FOR A.I.F.B.C. 8P - FLUID COMPRESSIBILITY BT - FLUID COEFFICIENT OF THERMAL EXPANSION **BV** - **VISCOSITY FUNCTION THERMAL PARAMETER** C - SOLUTE MASS FRACTION - SOLUTE CONCENTRATION AT THE END OF A WELL RISER, INITIAL VALUE C00 C11 - C11 THROUGH C33 ARE COEFFICIENTS IN THE CAPACITANCE MATRIX C12 C13 C21 C22

C23 C24 - C24,C34, AND C35 ARE COEFFICIENTS OF THE TRANSPORT TERMS C31 C32 C33 **C34** C35 CAIF - SOLUTE WASS FRACTION IN OUTER AQUIFER REGION **CC24** - CC24,CC34,CC35 ARE CONDUCTANCE COFACTOR ARRAYS **CC34** CC35 CCLBL - LABEL FOR SOLUTE MASS FRACTION IN A WELL FOR TEMPORAL PLOTS **U**JJ - CALCULATED SOLUTE HASS FRACTION IN A WELL CFLX - SOLUTE WASS FRACTION FOR INFLOW AT A SPECIFIED FLUX B.C. **CHAPRT - CHARACTER ARRAY FOR PRINTOUT** CHARC - 'C' CHARS - NUMERIC CHARACTERS THAT ILLUSTPATE THE CONTOUR-MAP ZONES CHKPTD - TRUE IF CHECK POINT DUMPS ARE DESIRED CI - NODAL CONNECTION INDEX FOR D4 NUMBERING SCHEME - CHARACTER REPRESENTATION OF IBC CIBC **CICALL - CHARACTER REPRESENTATION OF ICALL** - SOLUTE MASS FRACTION FOR INFLOW AT LEAKAGE B.C. CLBC - CHARACTER LINE FOR CONTOUR MAP CLINE - SOLUTE CONCENTRATION AT CELL FACE BETWEEN X(I-1) AND X(I) CHX - SOLUTE CONCENTRATION AT CELL FACE BETWEEN Y(J-1) AND Y(J) CHY - SOLUTE CONCENTRATION AT CELL FACE BETWEEN 2(K-1) AND 2(K) CHZ CNP - SOLUTE WASS FRACTION FOR SPECIFIED VALUE B.C. CNV - CONVERSION FACTOR - CONVERSION FACTOR FOR DENSITY (ENGLISH TO HETRIC) CNVD CHVDFI - CONVERSION FACTOR FOR DIFFUSIVITY (METRIC TO ENGLISH) CNVDI - CONVECTOR FOR DENSITY (METRIC TO ENGLISH) - CONVERSION FACTOR FOR ENERGY (ENGLISH TO METRIC) CHVE CHVEPI - CONVERSION FACTOR FOR PRESSURE ENERGY (METRIC TO ENGLISH) CHVFF - CONVERSION FACTOR FOR FLUID FLUX (ENGLISH TO HETRIC) CNVGZ - CONVERSION FACTOR FOR GRAVITY TIMES ELEVATION (ENGLISH TO METRIC) CHVIC - CONVERSION FACTOR FOR HEAT CAPACITY (ENGLISH TO HETRIC) CNVIICI - CONVERSION FACTOR FOR HEAT CAPACITY (HETRIC TO ENGLISH) CNVHF - CONVERSION FACTOR FOR HEAT FLUX (ENGLISH TO HETRIC) CNVINFI - CONVERSION FACTOR FOR HEAT FLUX (METRIC TO ENGLISH) CNVIII - CONVERISON FACTOR FOR THERMAL ENERGY (METRIC TO ENGLISH) CNVHTC - CONVERSION FACTOR FOR HEAT TRANSFER COEFFICIENT (ENGLISH TO METRIC) CNVHTI - CONVERSION FACTOR FOR HEAT TRANSFER COEFFICIENT (METRIC TO ENGLISH) CHVL - CONVERSION FACTOR FOR LENGTH (ENGLISH TO HETRIC) CNVL2 - CONVERSION FACTOR FOR LENGTH SQUARED (ENGLISH TO METRIC) CNVL21 - CONVERSION FACTOR FOR LENGTH SQUARED (METRIC TO ENGLISH)

CNVL3 - CONVERISON FACTOR FOR LENGTH CUBED (ENGLISH TO METRIC) CNVL31 - CONVERSION FACTOR FOR LENGTH CUBED (METRIC TO ENGLISH) CHVLI - CONVERSION FACTOR FOR LENGTH (METRIC TO ENGLISH) CHVHFI - CONVERSION FACTOR FOR MASS FLOW RATE (METRIC TO ENGLISH) CNVMI - CONVERSION FACTOR FOR MASS (METRIC TO ENGLISH) CNVP - CONVERSION FACTOR FOR PRESSURE (ENGLISH TO METRIC) CHYPI - CONVERSION FACTOR FOR PRESSURE (METRIC TO ENGLISH) CHVSF - CONVERSION FACTOR FOR SOLUTE FLUX (ENGLISH TO METRIC) CNVT1 - FIRST CONVERSION FACTOR FOR TEMPERATURE (ENGLISH TO HETRIC) CNVT11 - FIRST CONVERSION FACTOR FOR TEMPERATURE (METRIC TO ENGLISH) CNVT2 - SECOND CONVERSION FACTOR FOR TEMPERATURE (ENGLISH TO METRIC) CNVT2I - SECOND CONVERSION FACTOR FOR TEMPERATURE (HETRIC TO ENGLISH) CHVTC - CONVERSION FACTOR FOR THERMAL CONDUCTIVITY (ENGLISH TO HETRIC) CNVTCI - CONVERSION FACTOR FOR THERNAL CONDUCTIVITY (METRIC TO ENGLISH) CNVTM - CONVERSION FACTOR FOR TIME (DAYS TO SECONDS) CHVTHI - CONVERSION FACTOR FOR TIME (SECONDS TO DAYS) CHVUUL - CONVERSION FACTOR FOR FLUID CONDUCTANCE (METRIC TO ENGLISH) CNVVF - CONVERSION FACTOR FOR VOLUMETRIC FLOW RATE (ENGLISH TO METRIC) CNVVLI - CONVERSION FACTOR FOR VELOCITY (METRIC TO ENGLISH) CHVVSI - CONVERSION FACTOR FOR VISCOSITY (KG/M-S TO CP) COLS - NUMBER OF COLUMNS REQUIRED FOR CONTOUR MAPS COMOPT - TRUE IF COMPUTATION OF AN OPTIMUM OVER-RELAXATION FACTOR IS TO BE DONE **CONLBL - LABEL FOR SOLUTE CONCENTRATION PRINTOUT** CONVC - TRUE IF SOLUTE CALCULATION HAS CONVERGED FOR A GIVEN TIME STEP CONVP - TRUE IF PRESSURE CALCULATION HAS CONVERGED FOR A GIVEN TIME STEP CONVRG - TRUE IF WELL-RISER CALCULATION HAS CONVERGED CONVT - TRUE IF TEMPERATURE CALCULATION HAS CONVERGED FOR A GIVEN TIME STEP CON - SOLUTE MASS FRACTION OBSERVED VALUES IN A WELL CPAR - CHARACTER REPRESENTATION OF INPUT PARAMETER CPARE - INITIAL CHARACTER REPRESENTATION OF INPUT PARAMETER CPF - HEAT CAPACITY OF FLUID AT MINIMUM SOLUTE CONCENTRATION CPX - SOLUTE CONCENTRATION AT CELL FACE BETWEEN X(I) AND X(I+1) CPY - SOLUTE CONCENTRATION AT CELL FACE BETWEEN Y(J) AND Y(J+1) SOLUTE CONCENTRATION AT CELL FACE BETWEEN Z(K) AND Z(K+1) CPZ CSBC - SOLUTE MASS FRACTION FOR INFLOW AT A SPECIFIED PRESSURE B.C. CVIS - SOLUTE CONCENTRATION DATA FOR CONCENTRATION-VISCOSITY TABLE CWKT - SOLUTE MASS FRACTION AT WELL DATUM LEVEL CYLIND - TRUE IF A CYLINDRICAL COORDINATE SYSTEM IS BEING USED - DIAGONAL ELEMENTS OF THE SYSTEM EQUATION MATRIX n DAMWRC - DAMPING FACTOR FOR WELL RISER CALCULATION DASHES - '-----' DBKD - POROUS MEDIUM BULK DENSITY TIMES LINEAR EQUILIBRIUM SORPTION COEFFICIENT DC - CHANGE IN SOLUTE MASS FRACTION DCMAX - MAXIMUM CHANGE IN SOLUTE MASS FRACTION DCTAS - SPECIFIED MAXIMUM CHANGE IN SOLUTE MASS FRACTION FOR AUTOMATIC TIME STEP ADJUSTMENT

DONMAX - MAXIMUM CHANGE IN DENSITY DOV - CHANGE IN DEPENDENT VARIABLE FOR FLOW AT SPECIFIED VALUE B.C. DECLAN - SOLUTE DECAY FACTOR DEHIR - CHANGE IN ENTHALPY IN THE REGION **DELTIM - CURRENT TIME STEP** DELX - NODE SPACING IN X-DIRECTION DELY - NODE SPACING IN Y-DIRECTION - NODE SPACING IN Z-DIRECTION DELZ DEN - FLUID DENSITY DENO - FLUID DENSITY AT REFERENCE CONDITIONS DENC - COEFFICIENT FOR FLUID DENSITY VARIATION WITH SOLUTE MASS FRACTION DENCHC - DENSITY CHANGE DUE TO SOLUTE CONCENTRATON CHANGES DENCHT - DENSITY CHANGE DUE TO TEMPERATURE CHANGES DENFO - FLUID DENSITY AT MINIMUM SOLUTE WASS FRACTION DENF1 - FLUID DENSITY AT HAXIMUN SOLUTE HASS FRACTION DENFBC - FLUID DENSITY FOR INFLOW AT SPECIFIED FLUX B.C. DENGL - CHANGE IN PRESSURE ESTIMATE OVER WELL RISER LENGTH DENLBC - FLUID DENSITY FOR INFLOW AT LEAKAGE B.C. DENN - FLUID DENSITY AT TIME LEVEL N DENNP - FLUID DENSITY AT TIME LEVEL N+1 DENGAR - FLUID DENSITY IN OUTER AQUIFER REGION FOR A.I.F.B.C. DENP - COEFFICIENT FOR FLUID DENSITY VARIATION WITH PRESSURE DENT - COEFFICIENT FOR FLUID DENSITY VARIATION WITH TENPERATURE - FLUID DENSITY IN A WELL BORE DENW DENNKT - FLUID DENSITY IN A WELL BORE AT THE WELL DATUM DENWRK - FLUID DENSITY IN A WELL RISER AT LEVEL K DET - DETERMINANT OF THE WELL RISER EQUATIONS DFIR - CHANGE IN FLUID IN THE REGION DH - NOLECULAR DIFFUSIVITY OF THE SOLUTE DOTS - \*....\* DP - CHANGE IN PRESSURE DPHAX - NAXIMUM CHANGE IN PRESSURE DPTAS - SPECIFIED HAXINUM CHANGE IN PRESSURE FOR AUTOMATIC TIME STEP ADJUSTMENT DPUDT - DERIVATIVE OF THE TRANSIENT AQUIFER INFLUENCE FUNCTION WITH TIME OPWKT - CHANGE IN PRESSURE AT A WELL DATUM DOFDP - DERIVATIVE OF FLOW RATE WITH PRESSURE DOHBC - CHANGE IN HEAT FLOW FOR ANY B.C. DQHCDT - DERIVATIVE OF HEAT FLOW RATE WITH TEMPERATURE FOR HEAT CONDUCTION B.C. DONOP - DERIVATIVE OF HEAT FLOW WITH PRESSURE DOHOT - DERIVATIVE OF HEAT FLOW WITH TEMPERALURE DOINDP - DERIVATIVE OF WELL HEAT FLOW WITH PRESSURE DOIWDT - DERIVATIVE OF WELL HEAT FLOW WITH TEMPERATURE DOSBC - CHANGE IN SOLUTE FLOW RATE FOR ANY B.C. DQSDC - DERIVATIVE OF SOLUTE FLOW RATE WITH CONCENTRATION DOSDP - DERIVATIVE OF SOLUTE FLOW RATE WITH PRESSURE

DOSWDC - DERIVATIVE OF WELL SOLUTE FLOW WITH CONCENTRATION DOSWDP - DERIVATIVE OF WELL SOLUTE FLOW WITH PRESSURE DOWDP - DERIVATIVE OF WELL FLOW RATE WITH PRESSURE DOWDPL - DERIVATIVE OF WELL FLOW RATE AT A LAYER WITH PRESSURE DOWLYR - CHANGE IN WELL FLOW RATE AT A LAYER DSIR - CHANGE IN SOLUTE IN THE REGION - CHANGE IN TEMPERATURE DT DTADZW - DERIVATIVE OF AMBIENT TEMPERATURE AT A WELL RISER WITH DISTANCE IN THE Z-DIRECTION DTHAWR - THERMAL DIFFUSIVITY OF THE MEDIUM SURROUNDING A WELL RISER DTHIC - THERMAL DIFFUSIVITY OF THE EXTERNAL MEDIUM AT A HEAT CONDUCTION BOUNDARY DTIMMN - SPECIFIED MINIMUM TIME STEP FOR AUTOMATIC TIME STEP ADJUSTMENT DTIMMX - SPECIFIED MAXIMUM TIME STEP FOR AUTOMATIC TIME STEP ADJUSTMENT DTMAX - MAXIMUM CHANGE IN TEMPERATURE DTTAS - SPECIFIED NAXIMUM CHANGE IN TEMPERATURE FOR AUTOMATIC TIME STEP ADJUSTMENT. - X-COORDINATE SPACING FOR PLOT OF POROUS MEDIA ZONES DX - Y-COORDINATE SPACING FOR PLOT OF POROUS MEDIA ZONES DY DYY-- CHANGE IN DEPENDENT VARIABLE VECTOR FOR WELL RISER INTEGRATION DZ - CHANGE IN Z-COORDINATE VALUE DZNIN - SPECIFIED MINIMUM CHANGE IN Z-COORDINATE FOR WELL RISER INTEGRATION EEUNIT - TRUE IF ENGLISH ENGINEERING UNITS ARE BEING USED FOR DATA INPUT AND OUTPUT EH - FLUID ENTHALPY - FLUID ENTHALPY AT REFERENCE CONDITIONS EHO - ENTHALPY AT THE END OF A WELL RISER. INITIAL VALUE EHOO - FLUID ENTHALPY TABLE FOR DEVIATIONS FROM SATURATED CONDITIONS WITH PRESSURE AND TEMPERATURE EHDT EHIR - ENTHALPY IN THE REGION EHIRO - INITIAL ENTHALPY IN THE REGION EHIRN - ENTHALPY IN THE REGION AT TIME LEVEL N EHMX - ENTHALPY AT CELL FACE BETWEEN X(I-1) AND X(I) EHMY - ENTHALPY AT CELL FACE BETWEEN Y(J-1) AND Y(J) EHMZ - ENTHALPY AT CELL FACE BETWEEN Z(K-1) AND Z(K) - ENTHALPY AT CELL FACE BETWEEN X(I) AND X(I+1) EHPX - ENTHALPY AT CELL FACE BETWEEN Y(J) AND Y(J+1) EHPY - ENTHALPY AT CELL FACE BETWEEN Z(K) AND Z(K+1) EHPZ EHST - FLUID ENTHALPY TABLE AT SATURATION AS A FUNCTION OF TEMPERATURE EHWEND - ENTHALPY AT THE END OF THE WELL RISER EHWKT - ENTHALPY IN A WELL AT THE WELL DATUM EHWSUR - ENTHALPY IN A WELL AT THE LAND SURFACE - WELL-RISER PIPE ROUGHNESS DIVIDED BY PIPE INSIDE DIAMETER EOD - TOLERANCE FACTOR FOR TEMPERATURE CALCULATED FROM ENTHALPY AND PRESSURE EPS EPSFAC - TOLERANCE FACTOR FOR TWO-LINE SUCCESSIVE OVERRELAXATION CALCULATION EPSOMG - TOLERANCE FACTOR FOR CONVERGENCE OF OPTIMUM OVER-RELAXATION PARAMETER CALCULATION EPSSOR - TOLERANCE FOR CONVERGENCE OF TWO-LINE SUCCESSIVE-OVER-RELAXATION EQUATION SOLVER EPSWR - TOLERANCE FOR CONVERGENCE OF WELL RISER CALCULATION ERREX - TRUE IF PROGRAM ABORT DUE TO ERROR CONDITIONS ERREXE - TRUE IF PROGRAM ABORT DUE TO EXECUTION ERRORS

ERREXT - TRUE IF PROGRAM ABORT DUE TO INPUT ERRORS EVMAX - MAXIMUM EIGENVALUE ESTIMATE. EVMIN - MINIMUM EIGENVALUE ESTINATE: - 181 EX EXTRAP - EXTRAPOLATED RESULTS OF THE WELL-RISER INTEGRATION FIAIF - FACTOR FOR AQUIFER INFLUENCE FUNCTION B.C. F2AIF - FACTOR FOR AQUIFER INFLUENCE FUNCTION B.C. FCJ - FUNCTION FOR WELL RISER HEAT TRANSFER USING CARSLAW-JAEGER SOLUTION FUOP - FRACTIONAL CHANGE IN WELL-DATUM PRESSURE FDSMTH - FACTOR FOR SPATIAL DIFFERENCING METHOD FOTMTH - FACTOR FOR TEMPORAL DIFFERENCING METHOD FEPHL - FRICTIONAL HEAD LOSS IN WELL RISER - FLUID IN THE REGION FIR - INITIAL FLUID IN THE REGION FIRO FIRN - FLUID IN THE REGION AT TIME LEVEL N FIRVO - INITIAL VOLUMETRIC AMOUNT OF FLUID IN THE REGION FLOREV - TRUE IF FLOW REVERSAL OCCURS AT A GIVEN LEVEL IN A WELL - TRUE IF A WELL IS PRODUCING OR INJECTING AT AT LEAST ONE LEVEL FLOY - CHARACTER STRING FOR PRINT FORMAT FHT FPR3 - 4\*PI\*R\*\*3 FRAC - FRACTION OF CELL THAT IS FILLED FOR UNCONFINED FLOW FRACH - FRACTION OF CELL FILLED AT TIME LEVEL N FRACHP - FRACTION OF CELL FILLED AT TIME LEVEL N+1 FRESUR - TRUE IF A FREE SURFACE BOUNDARY IS ALLOWED FRFAC - FRICTION FACTOR FOR FLOW IN A WELL RISER PIPE FRFLM - FRICTION FACTOR IN WELL BORE AT BOUNDARY BETWEEN LEVEL L-1 AND L FRFLP - FRICTION FACTOR IN WELL BORE AT BOUNDARY BETWEEN LEVEL L AND L+1 - FUNCTION TABLE FOR INTERPOLATION FS FSCON - TRUE IF A FREE SURFACE RISES ABOVE THE TOP OF A CELL AND CONVERTS TO CONFINED CONDITIONS FSLOW - TRUE IF FREE SURFACE FALLS BELOW BOTTOM OF CELL FTDAIF - FACTOR FOR DIMENSIONLESS TIME FOR A.I.F.B.C. GCOSTH - COMPONENT OF GRAVITY ALONG A WELL RISER GRAV - GRAVITATIONAL CONSTANT GX - COMPONENT OF GRAVITY IN THE X-DIRECTION GY - COMPONENT OF GRAVITY IN THE Y-DIRECTION - CONPONENT OF GRAVITY IN THE Z-DIRECTION GZ HOPRNT - POTENTIONETRIC HEAD FOR PRINTOUT HEAT - TRUE IF HEAT TRANSPORT IS BEING SINULATED HICU - OVERALL HEAT TRANSFER COEFFICIENT FOR WELL RISER CALCULATON HITCWR - HEAT TRANSFER COEFFICIENT FROM THE FLUID TO A WELL RISER PIPE - ELEVATION OF THE WATER TABLE HUT THE UNDEFINED 'I' VARIABLES ARE POINTERS FOR THE LARGE VARIABLY PARTITIONED ARRAYS 112X - 12X' FORMAT 117 - ZONE INDEX - ZONE INDEX 122

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1BC - INDEX OF BOUNDARY CONDITION TYPE FOR A CELL **IBCMAP - INDEX OF BOUNDARY CONDITION TYPE FOR CONTOUR MAP** ICALL - INDEX OF CALL TO ARRAY-INPUT ROUTINE ICC - TRUE IF INITIAL CONDITION SOLUTE MASS FRACTION VALUES TO BE READ ICHWT - TRUE IF INITIAL CONDITION WATER-TABLE ELEVATION TO BE READ ICHYDP - TRUE IF INITIAL CONDITION OF HYDROSTATIC PRESSURE DISTRIBUTION ICMAX - INDEX IN X-DIRECTION OF CELL WITH MAXIMUM CHANGE IN SOLUTE CONCENTRATION ICON - INDEX OF CONNECTED NODES ICT - TRUE IF INITIAL CONDITION TEMPERATURE DISTRIBUTION TO BE READ ICXM - INDEX FOR VA ARRAY OF CONNECTING NODE IN NEGATIVE X-DIRECTION ICXP - INDEX FOR VA ARRAY OF CONNECTING NODE IN POSITIVE X-DIRECTION ICYN - INDEX FOR VA ARRAY OF CONNECTING NODE IN NEGATIVE Y-DIRECTION ICYP - INDEX FOR VA ARRAY OF CONNECTING NODE IN POSITIVE Y-DIRECTION ICZH - INDEX FOR VA ARRAY OF CONNECTING NODE IN NEGATIVE Z-DIRECTION ICZP - INDEX FOR VA ARRAY OF CONNECTING NODE IN POSITIVE Z-DIRECTION 104 - INDEX FOR D4 NODE RENUMBERING ID4CON - INDEX OF CONNECTED NODES FOR D4 NODE REMUMBERING ID4NO - INDEX OF NEW NODE NUMBERS FOR D4 REMUMBERING SCHEME IDIAG - INDEX OF DIAGONAL ELEMENTS IN THE A4 MATRIX IDIR - INDEX OF DIRECTION FOR TWO-LINE-SOR SOLVER IDNUM - IDENTIFICATION NUMBER FOR TEMPORAL PLOT **IDRCON - INDEX OF DIRECTION TO NEIGHBORING CONNECTED NODE** IE - INDEX OF ERROR NUMBER IE1 - INDEX LIMIT OF ERROR NUMBER IE2 - INDEX LIMIT OF ERROR NUMBER IENDIV - INDEX OF LAST LOCATION IN THE IVPA ARRAY IENDVV - INDEX OF LAST LOCATION IN THE VPA ARRAY - INDEX OF EQUATION BEING SOLVED 160 - INDEX OF ERROR NUMBER IER ' IERR - LOGICAL FLAGS FOR ERROR MESSAGES IF12 - 'F12' FORMAT IFACE - INDEX OF FACE OF APPLIED B.C. IFMT - INDEX OF FORMAT TYPE IFORM - CHARACTER STRING CONTAINING FORMAT STATEMENT IG12 - 'G12' FORMAT 1112 - '112' FORMAT 115 - '15' FORMAT ILAVEL - TOTAL LENGTH OF ALL LABELED COMMON BLOCKS ILBL - INDEX OF LABEL CHARACTERS FOR TEMPORAL PLOT ILH - INDEX OF ELEMENT IN EQUATION MATRIX FOR D4 SOLVER ILIVPA - LENGTH OF IVPA ARRAY AS DIMENSIONED 1LNXI = (LOG(XI))\*\*-1ILVPA - LENGTH OF VPA ARRAY AS DIMENSIONED IMAP1 - IMAP1. IMAP2. JMAP1. JMAP2. KMAP1. KMAP2 ARE NODE RANGES FOR MAP OUTPUT IMAP2

INOD	- INDEX OF HOB!FICATION FOR ARRAY INPUT
IMPPTC	~ INDEX FOR CONTOUR MAP OF DEPENDENT VARIABLE ARRAYS
INPQH	~ TRUE IF SEMI-IMPLICIT WELL FLOW CALCULATION IS DESIRED
INZONE	- TRUE IF A CELL IS CONTAINED WITHIN A DEFINED POROUS MEDIA ZONE
IPAGE	- PAGE NUMBER FOR CONTOUR MAP
IPAR	- INTEGER PARAMETER VALUE FOR ARRAY INPUT
IPAR1	- INITIAL INTEGER PARAMETER VALUE FOR ARRAY INPUT
IPLOT	~ INDEX ARRAY IDENTIFYING EXCLUDED CELLS FOR CONTOUR MAPS
IPHAX	- INDEX IN X-DIRECTION OF CELL WITH MAXIMUM CHANGE IN PRESSURE
IPHZ	- INDEX OF POROUS MEDIUM ZONE
[PHZ]	- INDEX OF FIRST ZONE FOR ZONE PLOT
IPHZ2	- INDEX OF LAST ZONE FOR ZONE PLOT
	- INDEX OF PRINTING PRESSURE, TEMPERATURE, OR SOLUTE CONCENTRAION ARRAYS
IROU	- INDEX OF EQUATION NUMBER FOR D4 SOLVER
ISIGN	
ISORD	
ISUN	- SUM OF I, J, AND K INDICES FORMING A DIAGONAL PLANE
ITINE	
ITHAX	
ITNO	- ITERATION NUMBER IN TWO-LINE SOR SOLVER
ITNOC	
	- ITERATIONS USED FOR PRESSURE CALCULATION BY TWO-LINE SOR SOLVER
ITNOT	
ITRN	- ITERATION COUNT FOR SEQUENCE OF P.T AND C EQUATION SOLUTION CYCLES
I TRN1	
	- ITERATION COUNT FOR WELLBORF CALCULATION
	- ITERATION COUNT FOR WELLBORE DENSITY CALCULATION
	- ITERATION COUNT FOR P,T OR P,C SOLUTION CYCLE
ITRNP	- ITERATION COUNT FOR P AND WELL BORE SOLUTION CYCLE
IUH	- INDEX OF EQUATION IN UPPER HALF OF NATRIX FOR D4 SOLVER
IVPA	- INTEGER VARIABLY PARTITIONED ARRAY
19	- INDEX OF CELL NUMBER IN X-DIRECTION FOR A WELL
INEL	- INDEX OF WELL NUMBER
JIZ	- ZONE INDEX
JZZ	- ZONE INDEX
JCHAX	
JHAP1	
JHAP2	
JPNAX	~ INDEX IN Y-DIRECTION OF CELL WITH MAXIMUM CHANGE IN PRESSURE
JTHAX	- INDEX IN Y-DIRECTION OF CELL WITH WAXIMUM CHANGE IN TEMPERATURE
JW	- INDEX OF CELL NUMBER IN Y-DIRECTION FOR A WELL
ĸĩz	- ZONE INDEX
K2Z	- ZONE INDEX
KARHC	
	- INDEX IN Z-DIRECTION OF CELL WITH MAXIMUM CHANGE IN SOLUTE CONCENTRATION
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KLBC - CONDUCTANCE FACTOR FOR LEAKAGE B.C. KHAP1 KMAP2 - WELL NUMBER INDEX FOR OBSERVED DATA KO KOAR - PERMEABILITY OF OUTER ADUIFER REGION FOR A.1.F.B.C. KPNAX - INDEX IN Z-DIRECTION OF CELL WITH MAXIMUM CHANGE IN PRESSURE KTHAWR - THERMAL CONDUCTIVITY OF AMBIENT MEDIUM AT WELL RISER KTHE - THERMAL CONDUCTIVITY OF FLUID KTHWR - THERMAL CONDUCTIVITY OF WELL RISER PIPE KTHX - THERMAL CONDUCTIVITY OF FLUID AND POROUS MEDIUM IN X-DIRECTION KTHXPM - THERMAL CONDUCTIVITY OF POROUS MEDIUM IN X-DIRECTION KTHY - THERMAL CONDUCTIVITY OF FLUID AND POROUS MEDIUM IN Y-DIRECTION KTHYPM - THERMAL CONDUCTIVITY OF POROUS MEDIUM IN Y-DIRECTION KTHZ - THERMAL CONDUCTIVITY OF FLUID AND POROUS MEDIUM IN Z-DIRECTION KTHZPH - THERHAL CONDUCTIVITY OF POROUS MEDIUM IN Z-DIRECTION KTMAX - INDEX IN 2-DIRECTION OF CELL WITH MAXIMUM CHANGE IN TEMPERATURE KXX - PERHEABILITY IN X-DIRECTION KYY - PERMEABILITY IN Y-DIRECTION - PERMEABILITY IN Z-DIRECTION KZZ -LABEL - CHARACTER STRING FOR LABEL LBLDIR - LABEL FOR OPTIMUM DIRECTION FOR TWO-LINE SOR SOLVER LBLED - LABEL FOR SYSTEM EQUATION NUMBER - LEFT SIDE BAND WIDTH OF A4 ARRAY FOR EACH EQUATION LBW **LCBOTW - INDEX OF BOTTOM LAYER OF COMPLETION OF A WELL** LCROSD - TRUE IF THE CROSS-DERIVATIVE DISPERSION COEFFICIENTS ARE TO BE LUMPED INTO THE DIAGONAL TERMS LCTOPW - INDEX OF TOP LAYER OF COMPLETION OF A WELL LDASH - '-----LDOTS - '..... LENAX - LENGTH OF THE X-AXIS FOR A CONTOUR MAP LENAY - LENGTH OF THE Y-AXIS FOR A CONTOUR MAP LENAZ - LENGTH OF THE Z-AXIS FOR A CONTOUR MAP LGREN - LOGARITHM OF THE REYNOLDS NUMBER LIMAGE - LINE IMAGE FOR DATA INPUT LIMIT - LABEL FOR WELL-BORE CALCULATION CONSTRAINTS LINE - CHARACTER STRING FOR TEMPORAL PLOT LINLIM - LIMIT TO THE NUMBER OF PRINTER LINES FOR TEMPORAL PLOT LOCAIF - INDEX FOR LOCATION OF THE AQUIFER INFLUENCE FUNCTION B.C. LPRNT - INDEX FOR INCLUSION OF A NODE IN AN ARRAY PRINTOUT - LOGARITHM OF DIMENSIONLESS TIME FOR A.I.F. LTD - CELL NUMBER H. MA - CELL NUMBER FOR ASSEMBLY MAIFC - CELL NUMBERS FOR AQUIFER INFLUENCE FUNCTION B.C. MAPPTC - INDEX FOR OUTPUT OF ZONED CONTOUR MAPS OF PRESSURE, TEMPERATURE, OR SOLUTE CONCENTRATION MAXIT - MAXIMUM NUMBER OF ITERATONS FOR TWO-LINE SOR SOLVER MAXITI - MAXIMUM NUMBER OF ITERATIONS ALLOWED FOR CALCULATION OF OPTIMUM OVER-RELAXATION FACTOR

HAXIT2 - HAXINUM NUMBER OF ITERATIONS ALLOWED FOR TWO-LINE SOR SOLUTION MAXITH - MAXIMUM NUMBER OF ITERATIONS ALLOWED FOR THE SOLUTION CYCLE OF THE THREE SYSTEM EQUATIONS MAXORD - MAXIMUM ORDER OF POLYNONIAL ALLOWED FOR INTEGRATION OF THE WELL-RISER EQUATIONS MAXPTS - MAXIMUM NUMBER OF SPATIAL STEPS ALLOWED FOR WELL-RISER CALCULATION NETH - INDEX FOR EXTRAPOLATION METHOD SELECTION FOR WELL-RISER CALCULATION - CELL NUMBERS FOR SPECIFIED FLUX B.C. MFBC **HFLBL - LABEL FOR HASS FRACTION OR SCALED HASS FRACTION** MICBC - CELL NUMBERS FOR HEAT CONDUCTION B.C. HIJKH - CELL NUMBER AT I.J.K-1 HIJKP - CELL NUHBER AT I.J.K+1 HIJHK - CELL NUMBER AT 1, J-1, K HIJHKH - CELL NUMBER AT I.J-1.K-1 HIJHKP - CELL NUMBER AT I, J-1, K+1 HIJPK - CELL NUMBER AT I.J+1.K HIJPKH - CELL NUMBER AT I.J+1.K-1 HIJPKP - CELL NUMBER AT I.J+1.K+1 MINJK - CELL NUMBER AT 1-1.J.K MINJKM - CELL NUMBER AT I-1.J.K-1 MIMJKP - CELL NUNBER AT I-1, J.K+1 HIMJHK - CELL NUMBER AT I-1.J-1.K HIHJPK - CELL NUMBER AT I-1.J+1.K HINUS - '-' MIPJK - CELL NUMBER AT I+1.J.K HIPJKH - CELL NUMBER AT I+1.J.K-1 MIPJKP - CELL NUMBER AT 1+1, J, K+1 MIPJHK - CELL NUMBER AT I+1.J-1.K MIPJPK - CELL NUMBER AT I+1, J+1, K - CELL NUMBER AT A WELL-DATION LEVEL HKT - CELL NUMBERS FOR LEAKAGE B.C. MLBC MOBIL - MOBILITY FOR AN AQUIFER LAYER AT A WELL BORE HS8C - CELL NUMBERS FOR SPECIFIED VALUE B.C. - CELL NUMBER AT TOP OF REGION HT MTJP1 - CELL NUMBER AT TOP OF REGION AT I.J+1 MXITQU - MAXIMUM NUMBER OF ITERATIONS ALLOWED FOR WELL FLOW RATE CALCULATION NAC - COUNT OF ACTIVE CELLS NAIFC - NUMBER OF AQUIFER INFLUENCE FUNCTION B.C. CELLS NCHARS - NUMBER OF CHARACTERS NCPR - NUMBER OF CHARACTERS TO BE PRINTED FOR ZONE PLOT - NUMBER OF DIMENSIONS FOR TABULAR INTERPOLATION NDIM NEHST - NUHBER OF ENTRIES IN THE SATURATED ENTHALPY VS. TEMPERATURE TABLE NFBC - NUMBER OF SPECIFIED FLUX B.C. CELLS NFC - COUNT OF SPECIFIED FLUX CELLS NGRIDX - COLUMN NUMBER CORRESPONDING TO X-GRID POINT NGRIDY - LINE NUMBER CORRESPONDING TO Y-GRID POINT NHC - COUNT OF HEAT CONDUCTION BOUNDARY CELLS

NHCBC - NUMBER OF HEAT CONDUCTION B.C. CELLS NHCN - NUMBER OF NODES EXTERNAL TO THE REGION FOR THE HEAT CONDUCTION B.C. CALCULATION NLBC - NUMBER OF LEAKAGE B.C. CELLS NLC - COUNT OF LEAKAGE B.C. CELLS • NMAPR. - NUMBER OF MAP RECORDS WRITTEN TO DISC NMPZON - NUMBER OF ZONES FOR CONTOUR MAPS NNOPPR - NUMBER OF NODES TO BE PRINTED FOR ARRAY OUTPUT NNPR - NUMBER OF NODES TO HAVE PRINTED VALUES - NUMBER OF OBSERVED DATA POINTS NO NOCA - NUMBER OF DATA POINTS FOR VISCOSITY VS. SOLUTE CONCENTRATION NOTY - NUMBER OF DATA POINTS FOR VISCOSITY VS. TEMPERATURE NOTVO - NUMBER OF DATA POINTS FOR VISCOSITY VS. TEMPERATURE AT MINIMUM SOLUTE CONCENTRATION NOTV1 - NUMBER OF DATA POINTS FOR VISCOSITY VS. TEMPERATURE AT MAXIMUM SOLUTE CONCENTRATION NPAGES - NUMBER OF PAGES FOR A CONTOUR MAP NPEHDT - NUMBER OF POINTS IN THE ENTHALPY DEVIATION TABLE ALONG THE PRESSURE COORDINATE NPMZ - NUMBER OF ZONES OF POROUS MEDIA PROPERTIES NPOSNS - NUMBER OF CHARACTER POSITIONS IN THE X-DIRECTION FOR A CONTOUR HAP ELEMENT NPTAIF - NUMBER OF POINTS IN THE TABLE OF TRANSIENT AQUIFER INFLUENCE FUNCTION VS. TIME NPTCBC - NUMBER OF SPECIFIED VALUE B.C. CELLS NPTSA4 - NUMBER OF POINTS IN THE A4 NATRIX FOR THE D4 SOLVER NPTSD4 - NUMBER OF POINTS FOR THE D4 SOLVER NPTSUH - NUMBER OF POINTS IN THE UPPER HALF OF THE EQUATION MATRIX FOR THE D4 SOLVER - NUMBER OF NODES IN THE R-DIRECTION NR. NRSTTP - NUMBER OF RESTART RECORD TIME PLANES WRITTEN TO DISC NSC - NUMBER OF SPECIFIED VALUE B.C. CELLS NSHUT - NUMBER OF WELLS SHUT IN NSTD4 - NUMBER OF STORAGE LOCATIONS REQUIRED BY THE D4 SOLVER NSTSOR - NUMBER OF STORAGE LOCATIONS REQUIRED BY THE TWO-LINE SOR SOLVER NTEHDT - NUMBER OF POINTS ALONG THE TEMPERATURE COORDINATE IN THE ENTHALPY DEVIATION TABLE NTHPTC - INDEX FOR SPECIFYING THAT EVERY NTH CALCULATED P, T OR C POINT IS TO BE PLOTTED NTHPTO - INDEX FOR SPECIFYING THAT EVERY NTH OBSERVED P, T OR C POINT IS TO BE PLOTTED NTSCHK - NUMBER OF TIME STEPS BETWEEN CHECKPOINT DUMPS NTSOPT - NUMBER OF TIME STEPS BETWEEN RECALCULATION OF THE OPTIMUM OVER-RELAXATION PARAMETER NVST - NUMBER OF POINTS IN THE TABLE OF GENERALIZED VISCOSITY VS. TEMPERATURE MMEL - NUMBER OF WELLS IN THE REGION - NUMBER OF NODES IN THE X-DIRECTION NX NXPR - NUMBER OF VALUES IN X-DIRECTION FOR ARRAY PRINTOUT - NUMBER OF NODES IN AN X-Y PLANE NXY NXYZ - NUMBER OF MODES IN THE SIMULATION REGION, NX\*NY\*NZ NY - NUMBER OF NODES IN THE Y-DIRECTION NYPR - NUMBER OF VALUES IN Y-DIRECTION FOR ARRAY PRINTOUT NZ - NUMBER OF NODES IN THE Z-DIRECTION NZPR - NUMBER OF VALUES IN Z-DIRECTION FOR ARRAY PRINTOUT NZIPHC - NUMBER OF NODES FOR THE INITIAL CONDITION TEMPERATURE PROFILE OUTSIDE THE REGION FOR HEAT CONDUCTION B.C. NZIPRO - NUMBER OF NODES FOR THE INITIAL CONDITION TEMPERATURE PROFILE IN THE Z-DIRECTION

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OCPLOT - TRUE IF PLOTS OF OBSERVED AND CALCULATED VALUES OF THE DEPENDENT VARIABLES ARE DESIRED OH - '0' OHEGA - OVER-RELAXATION FACTOR FOR THO-LINE SOR SOLVER OHOPT - OPTIMUM OVER-RELAXATION FACTOR FOR TWO-LINE SOR SOLVER ORENPR - ORIENTATION OF THE ARRAY PRINTOHITS, AREAL OR VERTICAL SLICE ORFR - ORIENTATION OF THE ARRAY PRINTOUIS FOR SATURATED FRACTION OF CELL P - PRESSURE PO - REFERENCE PRESSURE FOR DENSITY P00 - PRESSURE AT THE END OF A WELL RISER, INITIAL VALUE POH - REFERENCE PRESSURE FOR ENTHALPY **P1** - '.1' FORMAT - '.2' FORMAT P2 - '.4' FORMAT P4 - '.5' FORMAT P5 PAATH - ATHOSPHERIC PRESSURE IN ABSOLUTE UNITS PAEHDT - PRESSURE VALUES (ABSOLUTE) FOR ENTHALPY DEVIATION TABLE PAIF - PRESSURE OUTSIDE THE REGION FOR A.I.F.B.C. PAD - PARAMETER VALUE FOR ARRAY INPUT - PARAMETER VALUE FOR ARRAY INPUT PARA PARE - PARAMETER VALUE FOR ARRAY INPUT PARC - PARAMETER VALUE FOR ARRAY INPUT PCS - PRESSURE CALCULATED IN A WELL AT LAND SURFACE FOR TENPORAL PLOT PCW - PRESSURE CALCULATED AT A WELL DATUM FOR TEMPORAL PLOT PFSLOW - TRUE IF A MESSAGE IS TO BE PRINTED WHEN THE FREE SURFACE FALLS BELOW A CELL BOUNDARY PHILBC - POTENTIONETRIC HEAD ON THE OTHER SIDE OF AN ADUITARD FOR A LEAKAGE B.C. - '3.14159...' PΙ PINIT - PRESSURE INITIAL CONDITION AT A GIVEN ELEVATION PLBL '- LABEL FOR PRESSURE ARRAY PRIMTOUTS PLOTWC - TRUE IF TEMPORAL PLOTS OF SOLUTE CONCENTRATIONS IN WELLS ARE DESIRED PLOTWP - TRUE IF TEMPORAL PLOTS OF PRESSURES IN WELLS ARE DESIRED PLOTWT - TRUE IF TENPORAL PLOTS OF TENPERATURES IN WELLS ARE DESIRED PLTZON - TRUE IF A PLOT OF THE POROUS MEDIA ZONES IS DESIRED PLUS . . . PHCHV - POROUS MEDIUM COMPRESSIBILITY COEFFICIENT IN HEAT EQUATION PHCV - POROUS NEDIUM COMPRESSIBILITY COEFFICIENT IN FLOW EQUATION PHHV - POROUS MEDIUM THERMAL COEFFICIENT IN HEAT EQUATION, CURRENT VALUE PHP - PRESSURE FOR SPECIFIED VALUE B.C. PORDAR - POROSITY OF THE OUTER AQUIFER REGION FOR A.I.F.B.C. POROS - POROSITY POS - PRESSURE OBSERVED IN A WELL AT THE LAND SURFACE FOR TEMPORAL PLOT POSUP - TRUE IF THE VERTICAL AXIS OF THE CONTOUR MAP IS POSITIVE UPWARD POH - PRESSURES OBSERVED IN A WELL AT THE WELL DATUM FOR TEMPORAL PLOT PRBCF - TRUE IF PRINTOUT OF BOUNDARY CONDITION FLOW RATES IS DESIRED PRDV – TRUE IF PRINTOUTS OF DENSITY AND VISCOSITY FIELDS ARE DESIRED PRGFB - TRUE IF PRINTOUT OF GLOBAL FLOW BALANCE IS DESIRED

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PRIBCE - PRINTOUT INTERVAL FOR BOUNDARY CONDITION FLOW RATES PRIDY - PRINTOUT INTERVAL FOR DENSITY AND VISCOSITY FIELDS PRIGFB - PRINTOUT INTERVAL FOR GLOBAL FLOW BALANCE PRIKD - PRINTOUT INTERVAL FOR CONDUCTANCES AND DISPERSION COEFFICIENTS PRIMAP - PRINTOUT INTERVAL FOR ZONED CONTOUR MAPS PRIPTC - PRINTOUT INTERVAL FOR PRESSURE, TENPERATURE, AND SOLUTE MASS FRACTION FIELDS PRISLM - PRINTOUT INTERVAL FOR SOLUTION METHOD INFORMATION PRIVEL - PRINTOUT INTERVAL FOR VELOCITY FIELD PRIMEL - PRINTOUT INTERVAL FOR WELL INFORMATION PRKD - TRUE IF PRINTOUTS OF CONDUCTANCES AND DISPERSION COEFFICIENTS ARE DESIRED PRPTC - TRUE IF PRINTOUTS OF DEPENDENT VARIABLES P.T OR C ARE DESIRED PRSLM - TRUE IF PRINTOUT OF SOLUTION METHOD INFORMATION IS DESIRED PRTBC - TRUE IF PRINTOUT OF BOUNDARY CONDITION FLOW RATES IS DESIRED PRTCCM - TRUE IF PRINTOUT OF MESSAGE OF FREE-SURFACE B.C., BECOMING CONFINED IS DESIRED PRICHR - CHARACTER STRING FOR PRINTOUT OF CONTOUR MAPS PRTDY - TRUE IF PRINTOUT OF DENSITY AND VISCOSITY FIELDS IS DESIRED PRTFP - TRUE IF PRINTOUT OF FLUID PARAMETERS IS DESIRED PRTIC - TRUE IF PRINTOUT OF INITIAL CONDITIONS IS DESIRED PRTMPD - TRUE IF PRINTOUT OF CONTOUR MAP PARAMETERS IS DESIRED PRTPMP - TRUE IF PRINTOUT OF POROUS MEDIA PROPERTIES IS DESIRED PRTRE - TRUE IF PRINTOUT OF READ ECHO FOR ARRAY INPUT IS DESIRED PRTSLM - TRUE IF PRINTOUT OF SOLUTION METHOD PARAMETERS IS DESIRED PRIVEL - TRUE IF PRINTOUT OF WELL BORE INFORMATION IS DESIRED PRVEL - TRUE IF PRINTOUT OF VELOCITY FIELD IS DESIRED PRWEL - TRUE OF PRINTOUT OF WELL BORE INFORMATION IS DESIRED PSBC - PRESSURE FOR SPECIFIED VALUE B.C. PSLBL - LABEL FOR WELL PRESSURE AT LAND SURFACE FOR TEMPORAL PLOTS PSHAX - MAXIMUM PRESSURE AT LAND SURFACE FOR SCALING TEMPORAL PLOTS PSHIN - MINIMUM PRESSURE AT LAND SURFACE FOR SCALING TEMPORAL PLOTS PU - TRANSIENT AQUIFER INFLUENCE FUNCTION DIMENSIONLESS PRESSURE RESPONSE TO A UNIT DIMENSIONLESS WITHDRAWAL FLOW-RATE CHAMGE - PORE VOLUME, CURRENT VALUE PV -- PORE VOLUME WEIGHTED SORPTION PARAMETER. CURRENT VALUE PVK PWCELL - PRESSURE IN THE CELL AT THE UPPERMOST COMPLETION LEVEL OF A WELL PWKT - PRESSURE IN A WELL AT THE WELL DATUM PWKTS - SPECIFIED LIMITING PRESSURE IN A WELL AT THE WELL DATUM PWLBL - LABEL FOR WELL PRESSURE AT WELL DATUM FOR TEMPORAL PLOTS PWMAX - MAXIHUM PRESSURE AT WELL DATUM FOR SCALING TEMPORAL PLOTS PHMIN - MINIMUM PRESSURE AT WELL DATUM FOR SCALING TEMPORAL PLOT PWREND - PRESSURE AT THE END OF A WELL RISER PIPE PWRK - PRESSURE IN A WELL RISER AT A GIVEN LEVEL PWSUR - PRESSURE IN A WELL AT THE LAND SURFACE PWSURS - SPECIFIED LIMITING PRESSURE IN A WELL AT THE LAND SURFACE ODVSBC - FLOW RATE FOR THE DEPENDENT VARIABLE AT A SECIFIED VALUE B.C. CELL QFAC - BI-LINEAR INTERPOLATION FACTOR FOR CONTOUR MAPS OFAIF - VOLUMETRIC FLOW RATE FOR A.I.F.B.C.

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OFBC - FLUID FLOW RATE AT ANY B.C. CELL OFBCV - VOLUMETRIC FLUID FLOW RATE FOR SPECIFIED FLUX B.C. - FLUID FLUX IN X-DIRECTION OFFX **ÖFFY** - FLUID FLUX IN Y-DIRECTION ÓFFZ - FLUID FLUX IN Z-DIRECTION OFLBC - FLUID WASS FLOW RATE FOR LEAKAGE B.C. OFSBC - VOLUMETRIC FLUID FLOW RATE FOR SPECIFIED VALUE B.C. OHAIF - HEAT FLOW RATE FOR A.I.F.B.C. OHBC ... - HEAT FLOW RATE AT ANY B.C. CELL OHCBC - HEAT FLOW RATE FOR HEAT CONDUCTION B.C. OHFAC - HEAT FLOW RATE FACTOR FOR WELL RISER OHFBC - HEAT FLOW RATE FOR SPECIFIED FLUX B.C. **OHFX** - SPECIFIED HEAT FLUX IN THE X-DIRECTION **OHFY** - SPECIFIED HEAT FLUX IN THE Y-DIRECTION **MIFZ** - SPECIFIED HEAT FLUX IN THE Z-DIRECTION QHLBC - HEAT FLOW RATE AT LEAKAGE B.C. QHLYR - HEAT FLOW RATE FROM A WELL TO AN AQUIFER LAYER OHSBC - HEAT FLOW RATE AT A SPECIFIED VALUE B.C. OHSBC - HEAT FLOW RATE AT A SPECIFIED PRESSURE B.C. OHU -- HEAT FLOW RATE FROM A WELL TO THE ADUIFER - HEAT FLOW RATE IN A WELL RISER AT A GIVEN LEVEL DINURK OL IN - LIMITING FLOW RATE FOR RIVER LEAKAGE ON . - FLOW RATE AT TIME LEVEL N - FLOW RATE AT TIME LEVEL N+1 ONP OSAIF - SOLUTE FLOW RATE AT AN A.I.F.B.C. - SOLUTE FLOW RATE AT ANY B.C. CELL OSBC **OSFBC** - SOLUTE FLOW RATE AT A SPECIFIED FLUX B.C. **ÓSFX** - SPECIFIED SOLUTE FLUX IN THE X-DIRECTION **OSFY** - SPECIFIED SOLUTE FLUX IN THE Y-DIRECTION OSFZ - SPECIFIED SOLUTE FLUX IN THE Z-DIRECTION **OSLBC - SOLUTE FLOW RATE AT A LEAKAGE B.C. QSLYR - SOLUTE FLOW RATE FROM A WELL TO AN AQUIFER LAYER OSSBC - SOLUTE FLOW RATE FOR A SPECIFIED VALUE B.C. ÚS₩** - SOLUTE FLOW RATE FROM A WELL TO THE AQUIFER OHAV - AVERAGE FLOW RATE FROM A WELL TO AN AQUIFER LAYER OVER A TIME STEP OHLYR - FLUID FLOW RATE FROM A HELL TO AN AQUIFER LAYER ONH. - FLUID HASS FLOW RATE FROM A WELL TO THE AQUIFER - FLUID FLOW RATE FROM A WELL TO THE AQUIFER AT TIME LEVEL N OWN OHNP - FLUID FLOW RATE OF A WELL AT TIME LEVEL N+1 - FLUID FLOW RATE IN A WELL RISER OHR OWV - VOLUMETRIC FLUID FLOW RATE FROM A WELL TO THE AQUIFER RBW - RIGHT-HAND BAND WIDTH FOR EACH EQUATION OF THE A4 MATRIX RCPPH - HEAT CAPACITY OF THE POROUS NEDIUM PER UNIT VOLUMF RDAIF - TRUE IF AQUIFER INFLUENCE B.C. INFORMATION IS TO BE READ RDCALC - TRUE IF CALCULATION INFORMATION IS TO BE READ

RDECHO - TRUE IF A READ-ECHO FILE IS TO BE WRITTEN RDFLXH - TRUE IF HEAT FLUX DATA ARE TO BE READ RDFLXO - TRUE IF FLUID FLUX DATA ARE TO BE READ RDFLXS - TRUE IF SOLUTE FLUX DATA ARE TO BE READ RDLBC - TRUE IF LEAKAGE B.C. DATA ARE TO BE READ RDMPDT - TRUE IF CONTOUR MAP INFORMATION IS TO BE READ ROPLTP - TRUE IF INFORMATION FOR PRESSURE PLOTS IS TO BE READ RDSCBC - TRUE IF SPECIFIED SOLUTE CONCENTRATION B.C. DATA ARE TO BE READ RDSPBC - TRUE IF SPECIFIED PRESSURE B.C. DATA ARE TO BE READ ROSTBC - TRUE IF SPECIFIED TEMPERATURE B.C. DATA ARE TO BE READ RDVAIF - TRUE IF A.I.F.B.C. GEOMETRIC FACTORS ARE TO BE READ ROWDEF - TRUE IF WELL DEFINITION INFORMATION IS TO BE READ ROWFLO - TRUE IF WELL FLOW RATE DATA ARE TO BE READ RDWHD - TRUE IF WELL-HEAD DATA AT LAND SURFACE ARE TO BE READ REN - REYNOLDS NUMBER FOR FLOW IN A WELL RISER OR WELL BORE **RESTRT - TRUE IF THIS IS A RESTART RUN** - RIGHT-HAND-SIDE VECTOR FOR THE FLOW EQUATION RF RH - RIGHT-HAND-SIDE VECTOR FOR THE HEAT EQUATION RH1 - RIGHT-HAND-SIDE VECTOR AUGMENTED WITH CROSS-DISPERSIVE FLUX TERMS RHS - RIGHT-HAND-SIDE VECTOR FOR THE SYSTEM EQUATIONS RHSSBC - RIGHT-HAND-SIDE VECTOR FOR THE SPECIFIED VALUE B.C. NODES RHSW - RIGHT-HAND-SIDE VECTOR FOR THE WELL-BORE NODES FOR CYLINDRICAL COORDINATE SYSTEM RIDAR - RADIUS OF THE APPROXIMATE BOUNDARY BETWEEN THE INNER AND OUTER AQUIFER REGIONS FOR A.I.F.B.C. RM - RADII OF CELL BOUNDARIES IN THE R-DIRECTION FOR THE CYLINDRICAL COORDINATE SYSTEM RORW2 - EQUIVALENT CELL RADIUS DIVIDED BY THE WELL RADIUS, QUANTITY SQUARED RPRN - '(' - RIGHT-HAND-SIDE VECTOR FOR THE SOLUTE EQUATION RS RS1 - RIGHT-HAND-SIDE VECTOR AUGMENTED WITH CROSS-DISPERSIVE FLUX TERMS SAVLDO - TRUE IF ONLY THE LAST RESTART OR CHECKPOINT DUMP IS TO BE SAVED SCALMF - TRUE IF A SCALED MASS FRACTION IS TO BE USED FOR INPUT AND OUTPUT SDECAY - AMOUNT OF SOLUTE DECAYED DURING A TIME STEP SHRES - RESIDUAL ERROR IN THE HEAT EQUATION FOR CURRENT TIME STEP SHRESF - FRACTIONAL RESIDUAL ERROR IN THE HEAT EQUATION FOR THE CURENT TIME STEP SIR - SOLUTE IN THE REGION SIRO - INITIAL SOLUTE IN THE REGION SIRN - SOLUTE IN THE REGION AT TIME LEVEL N SLMETH - INDEX FOR SELECTION OF EQUATION SOLUTION HETHOD SMCALC - TRUE IF AN ABOREVIATED CAPACITANCE-COEFFICIENT CALCULATION IS TO BE DONE SOLUTE - TRUE IF SOLUTE TRANSPORT IS BEING SIMULATED SOLVE - TRUE IF THE TWO-LINE-SOR SOLVER IS TO SOLVE THE EQUATIONS SPRAD - SPECTRAL RADIUS OF THE EQUATION COEFFICIENT MATRIX SRES - RESIDUAL ERROR IN THE FLOW EQUATION FOR THE CURRENT TIME STEP SRESF - FRACTIONAL RESIDUAL ERROR IN THE FLOW EQUATION FOR THE CURRENT TIME STEP SSRES - RESIDUAL ERROR IN THE SOLUTE EQUATION FOR THE CURRENT TIME STEP SSRESF - FRACTIONAL RESIDUAL ERROR IN THE SOLUTE EQUATION FOR THE CURRENT TIME STEP

ين 8

STOTFI - FLUID INPUT TO THE REGION DURING THE CURRENT TIME STEP STOTFP - FLUID PRODUCED FROM THE REGION DURING THE CURRENT TIME STEP STOTHI - HEAT INPUT TO THE REGION DURING THE CURRENT TIME STEP STOTHP - HEAT PRODUCED FROM THE REGION DURING THE CURRENT TIME STEP STOTSI - SOLUTE INPUT TO THE REGION DURING THE CURRENT TIME STEP STOTSP - SOLUTE PRODUCED FROM THE REGION DURING THE CURRENT TIME STEP SUMMOB - SUM OF ALL WELL MOBILITIES FOR A GIVEN WELL SUNWI - SUN OF ALL WELL INDICES FOR A GIVEN WELL - TRUE IF A SPECIFIED VALUE B.C. IS AT THIS CELL SVBC SXX - FLUID MASS FLOW RATE IN THE X-DIRECTION - FLUID MASS FLOW RATE IN X-DIRECTION BETWEEN CELL AT X(I-1) AND X(1) SXXM - FLUID MASS FLOW RATE IN X-DIRECTION BETWEEN CELL AT X(1) AND X(1+1) SXXP SYNID - CHARACTER SYMBOL FOR CONTOUR MAP BOUNDARIES AND NODES - FLUID WASS FLOW RATE IN THE Y-DIRECTION SYY SYYH - FLUID MASS FLOW RATE IN Y-DIRECTION BETWEEN CELL AT Y(J-1) AND Y(J) SYYP - FLUID WASS FLOW RATE IN Y-DIRECTION BETWEEN CELL AT Y(J) AND Y(J+1) SZZ - FLUID WASS FLOW RATE IN THE Z-DIRECTION SZZH - FLUID MASS FLOW RATE IN 2-DIRECTION BETWEEN CELL AT Z(K-1) AND Z(K) - FLUID MASS FLOW RATE IN Z-DIRECTION BETWEEN CELL AT Z(K) AND Z(K+1) SZZP - FLUID MASS FLOW RATE UP THE WELL BORE SZZW - TEMPERATURE T TO - REFERENCE TEMPERATURE FOR DENSITY TO. - REFERENCE TEMPERATURE AT WHICH DENSITY DATA ARE GIVEN T00 - TEMPERATURE AT THE END OF A WELL RISER, INITIAL VALUE - REFERENCE TEMPERATURE FOR ENTHALPY AND VISCOSITY TOH TABUR - TEMPERATURE IN THE AMBIENT MEDIUM AT THE BOTTOM OF A WELL RISER TAIF - TEMPERATURE IN THE OUTER AQUIFER REGION FOR A.I.F.B.C. TANBK - TENPERATURE IN THE ANBIENT NEDIUM AT A GIVEN LEVEL ALONG THE WELL RISER TATWR - TEMPERATURE IN THE AMBIENT MEDIUM AT THE TOP OF A WELL RISER TC - TIME VALUES FOR CALCULATED VARIABLES AT A WELL TCS - TEMPERATURE CALCULATED IN A WELL AT LAND SURFACE FOR TEMPORAL PLOT - TEMPERATURE CALCULATED AT A WELL DATUM FOR TEMPORAL PLOT TCH TOEHIR - CUMULATIVE CHANGE IN ENTHALPY IN THE REGION TOFIR - CUMULATIVE CHANGE IN FLUID IN THE REGION TDSIR - CUMULATIVE CHANGE IN SOLUTE IN THE REGION - DISPERSIVE CONDUCTANCE IN X-DIRECTION TDX TDXY - CROSS-DISPERSIVE CONDUCTANCE TOXZ - CROSS-DISPERSIVE CONDUCTANCE TOY - DISPERSIVE CONDUCTANCE IN Y-DIRECTION - CROSS-DISPERSIVE CONDUCTANCE TOYX TDYZ - CROSS-DISPERSIVE CONDUCTANCE TDZ - DISPERSIVE CONDUCTANCE IN Z-DIRECTION - CROSS-DISPERSIVE CONDUCTANCE TOZX - CROSS-DISPERSIVE CONDUCTANCE TOZY TEHDT - TEMPERATURE VALUES FOR THE TABLE OF ENTHALPY DEVIATION

TEHST - TEMPERATURE VALUES FOR THE TABLE OF ENTHALPY AT SATURATION TFLX - TEMPERATURE FOR INFLOW AT A SPECIFIED FLUX B.C. TFRES - CUMULATIVE RESIDUAL ERROR IN THE FLOW EQUATION TFRESF - FRACTIONAL CUMULATIVE RESIDUAL ERROR IN THE FLOW EQUATION TFW - CONDUCTANCE IN WELL BORE FOR CYLINORICAL SYSTEM TFX - FLUID CONDUCTANCE IN X-DIRECTION - FLUID CONDUCTANCE IN Y-DIRECTION TFY - FLUID CONDUCTANCE IN Z-DIRECTION TFZ THCBC - TEMPERATURE PROFILE FOR FIRST PROBLEM FOR HEAT CONDUCTION B.C. THETXZ - ANGLE BETWEEN X-AXIS AND GRAVITATIONAL VECTOR THETYZ - ANGLE BETWEEN Y-AXIS AND GRAVITATIONAL VECTOR THETZZ - ANGLE BETWEEN Z-AXIS AND GRAVITATIONAL VECTOR THRES - CUMULATIVE RESIDUAL ERROR IN THE HEAT EQUATION THRESF - FRACTIONAL CUMULATIVE RESIDUAL ERROR IN THE HEAT EQUATION THRU - TRUE IF THIS IS THE END OF THE SIMULATION - THERMAL CONDUCTANCE IN X-DIRECTION THX THXY - CROSS-DISPERSIVE THERMAL CONDUCTANCE THXZ - CROSS-DISPERSIVE THERMAL CONDUCTANCE THY - THERMAL CONDUCTANCE IN Y-DIRECTION - CROSS-DISPERSIVE THERMAL CONDUCTANCE THYX THYZ - CROSS-DISPERSIVE THERMAL CONDUCTANCE THZ - THERMAL CONDUCTANCE IN Z-DIRECTION THZX - CROSS-DISPERSIVE THERMAL CONDUCTANCE THZY - CROSS-DISPERSIVE THERMAL CONDUCTANCE TILT - TRUE IF THE COORDINATE SYSTEM IS TILTED SO THAT THE GRAVITATIONAL VECTOR DOES NOT POINT IN THE NEGATIVE 2-DIRECTION TIMCHG - TIME AT WHICH NEW TRANSIENT DATA ARE TO BE READ TIMDAY - TIME IN DAYS TIME - TIME VALUE TIMED - DIMENSIONLESS TIME TIMEDN - DIMENSIONLESS TIME AT TIME LEVEL N TIMRST - TIME AT WHICH A RESTART RECORD IS TO BE WRITTEN TITLE - TITLE FOR THE SIMULATION RUN TITLEO - TITLE FOR THE ORIGINAL SIMULATION THAT IS BEING RESTARTED TLBC - TEMPERATURE AT THE OTHER SIDE OF AN CONFINING LAYER FOR A LEAKAGE B.C. TLBL - LABEL FOR TEMPERATURE FOR ARRAY PRINTOUTS TM1 - CONDUCTANCE COEFFICIENT VECTOR FOR TWO-LINE SOR SOLVER - CONDUCTANCE COEFFICIENT VECTOR FOR TWO-LINE SOR SOLVER TH2 THLBL - LABEL FOR TIME FOR TEMPORAL PLOTS TNP - TEMPERATURE FOR SPECIFIED VALUE B.C. - TIME VALUES FOR OBSERVED VARIABLES IN A WELL FOR TEMPORAL PLOTS TO. TOL - TOLERANCE FOR AN ITERATIVE CALCULATION TOLULN - TOLERANCE ON DENSITY CHANGES FOR CONVERGENCE OF THE SOLUTION CYCLE OF THE THREE SYSTEM FOUNTIONS TOLDNC - TOLERANCE ON DENSITY CHANGES DUE TO SOLUTE CONCENTRATION CHANGES FOR THE SOLUTION CYCLE OF THE FLOW AND SOLUTE EQUATIONS TOLONT - TOLERANCE ON DENSITY CHANGES DUE TO TEMPERATURE CHANGES FOR THE SOLUTION CYCLE OF THE FLOW AND HEAT EQUATIONS TOLDPW - TOLERANCE ON PRESSURE CHANGES FOR THE WELL BORE CALCULATION

TOLFPW - TOLERANCE ON FRACTIONAL PRESSURE CHANGES FOR THE WELL-BORE CALCULATION TOLOW - TOLERANCE ON FLOW RATE CHANGES FOR THE WELL-BORE CALCULATION ZOT - TEMPERATURES OBSERVED IN A WELL AT THE LAND SURFACE FOR TEMPORAL PLOT TOTFI - CUMULATIVE FLUID INPUT TO THE REGION TOTEP - CUMULATIVE FLUID PRODUCED FROM THE REGION TOTHE - CUMULATIVE HEAT INPUT TO THE REGION TOTHP - CUMULATIVE HEAT PRODUCED FROM THE REGION TOTSI - CUMULATIVE SOLUTE INPUT TO THE REGION TOTSP - CUMULATIVE SOLUTE PRODUCED FROM THE REGION TOTWFI - CUMULATIVE FLUID INJECTED BY THE WELLS TOTHEP - CUMULATIVE FLUID PRODUCED BY THE WELLS TOTWHE - CUMULATIVE HEAT INJECTED BY THE WELLS TOTWHP - CUMULATIVE HEAT PRODUCED BY THE WELLS TOTWSI - CUMULATIVE SOLUTE INJECTED BY THE WELLS TOTWSP - CUMULATIVE SOLUTE PRODUCED BY THE WELLS - TENPERATURES OBSERVED IN A WELL AT THE WELL DATUH FOR TENPORAL PLOT TON TP1 - CONDUCTANCE COEFFICIENT VECTOR FOR TWO-LINE SOR SOLVER TP2 - CONDUCTANCE COEFFICIENT VECTOR FOR TWO-LINE SOR SOLVER TPHCBC - TEMPERATURE PROFILE FOR SECOND PROBLEM FOR HEAT CONDUCTION B.C. TOFAIF - CUMULATIVE FLUID INPUT FROM A.I.F.B.C. TOFFBC - CUMULATIVE FLUID INPUT FROM SPECIFIED FLUX B.C. TOFINJ - TOTAL FLUID INJECTION RATE FOR ALL WELLS TOFLEC - CUMULATIVE FLUID INPUT FROM LEAKAGE B.C. TOFPRO - TOTAL FLUID PRODUCTION RATE FROM ALL WELLS TOFSBC - CUMULATIVE FLUID INPUT FROM SPECIFIED VALUE B.C. TOHAIF - CUMULATIVE HEAT INPUT FROM A.I.F.B.C. TOHFBC - CUMULATIVE HEAT INPUT FROM SPECIFIED FLUX B.C. TOHHBC - CUMULATIVE HEAT INPUT FROM HEAT CONDUCTION B.C. TOHINJ - TOTAL HEAT INJECTION RATE FROM ALL WELLS TOHLBC - CUMULATIVE HEAT INPUT FROM LEAKAGE B.C. TOHPRO - TOTAL HEAT PRODUCTION RATE FROM ALL WELLS TOHSBC - CUMULATIVE NEAT INPUT FROM SPECIFIED VALUE B.C. TOSAIF - CUMULATIVE SOLUTE INPUT FROM A.I.F.B.C. TOSFBC - CUMULATIVE SOLUTE INPUT FROM SPECIFIED FLUX B.C. TOSINJ - TOTAL SOLUTE INJECTION RATE FROM ALL WELLS TOSLBC - CUMULATIVE SOLUTE INPUT FROM LEAKAGE B.C. TOSPRO - TOTAL SOLUTE PRODUCTION RATE FROM ALL WELLS TOSSBC - CUMULATIVE SOLUTE INPUT FROM SPECIFIED VALUE B.C. TRVIS - REFERENCE TEMPERATURE AT WHICH VISCOSITY DATA ARE GIVEN TSBC - TEMPERATURE FOR INFLOW AT A SPECIFIED PRESSURE B.C. TSLBL - LABEL FOR WELL TEMPERATURE AT LAND SURFACE FOR TEMPORAL PLOTS TSHAX - MAXINUM TEMPERATURE AT LAND SURFACE FOR SCALING TEMPORAL PLOTS TSMIN - MININUM TEMPERATURE AT LAND SURFACE FOR SCALING TEMPORAL PLOTS TSRES - CUMULATIVE RESIDUAL ERROR IN THE SOLUTE EQUATION **TSRESF - FRACTIONAL CUMULATIVE ERROR IN THE SOLUTE EQUATION** 

TSX - SOLUTE CONDUCTANCE IN X-DIRECTION TSXY - CROSS-DISPERSIVE SOLUTE CONDUCTANCE TSXZ - CROSS-DISPERSIVE SOLUTE: CONDUCTANCE TSY - SOLUTE CONDUCTANCE IN Y-DIRECTION TSYX - CROSS-DISPERSIVE SOLUTE CONDUCTANCE - CROSS-DISPERSIVE SOLUTE CONDUCTANCE TSYZ TSZ - SOLUTE CONDUCTANCE IN 2-DIRECTION TSZX - CROSS-DISPERSIVE SOLUTE CONDUCTANCE **TSZY** - CROSS-DISPERSIVE SOLUTE CONDUCTANCE TVD - TEMPERATURE VS. DEPTH VALUES FOR INITIAL CONDITION TYFO - TEMPERATURE VALUES FOR VISCOSITY VS. TEMPERATURE DATA AT MINIMUM SOLUTE CONCENTRATION TVF1 - TEMPERATURE VALUES FOR VISCOSITY VS. TEMPERATURE DATA AT MAXIMUM SOLUTE CONCENTRATION TVZHC - TEMPERATURE VS. DISTANCE OUTWARD FROM HEAT CONDUCTION B.C. FOR INITIAL CONDITION TWKT - TEMPERATURE IN A WELL AT THE WELL DATUM TWLBL - LABEL FOR WELL TEMPERATURE AT WELL DATUM FOR TEMPORAL PLOTS TWMAX - MAXIMUM TEMPERATURE AT WELL DATUM FOR SCALING TEMPORAL PLOTS TIMIN - NINIMUN TEMPERATURE AT WELL DATUM FOR SCALING TEMPORAL PLOTS TWOP1 - '6.28...' TWREND - TEMPERATURE AT THE END OF A WELL RISER PIPE TWRK - TEMPERATURE IN A WELL RISER AT A GIVEN LEVEL TWSRKT - TEMPERATURE AT LAND SURFACE OR AT WELL DATUM TWSUR - TEMPERATURE IN A WELL AT THE LAND SURFACE - FLUID CONDUCTANCE FACTOR IN THE X-DIRECTION TX. TY - FLUID CONDUCTANCE FACTOR IN THE Y-DIRECTION - FLUID CONDUCTANCE FACTOR IN THE Z-DIRECTION 1Z THE UNDEFINED 'U' VARIABLES ARE USED FOR TEMPORARY STORAGE UCROSC - CROSS-DISPERSIVE SOLUTE FLUX UCROST - CROSS-DISPERSIVE HEAT FLUX UNIGRX - TRUE IF A UNIFORM GRID IN THE X-DIRECTION IS DESIRED UNIGRY - TRUE IF A UNIFORM GRID IN THE Y-DIRECTION IS DESIRED UNIGRZ - TRUE IF A UNIFORM GRID IN THE Z-DIRECTION IS DESIRED UNITEP - UNIT FOR ENERGY PER UNIT MASS UNITH - UNIT FOR HEAT UNITHF - UNIT FOR HEAT FLOW RATE UNITE - UNIT FOR LENGTH UNITH - UNIT FOR MASS UNITP - UNIT FOR PRESSURE UNITE - UNIT FOR TEMPERATURE UNITTH - UNIT FOR TIME UNITVS - UNIT FOR VISCOSITY - SYSTEM EQUATION COEFFICIENT MATRIX VA VAIFC - GEOMETRIC FACTOR FOR A.I.F.B.C. VAR - VARIABLE FOR ARRAY INPUT VASBC - SYSTEM EQUATION COEFFICIENT MATRIX FOR NODES WITH SPECIFIED B.C. - SYSTEM EQUATION COEFFICIENT MATRIX FOR NODES ALONG WELL BORE IN CYLINDRICAL COORDINATES

VUATA - VISCOSITY DATA VELWRK - VELOCITY OF FLOW IN A WELL RISER AT A GIVEN LEVEL VIS - VISCOSITY VISCTR - VISCOSITY VS. SOLUTE CONCENTRATION DATA AT REFERENCE TEMPERATURE VISLBC - VISCOSITY AT OTHER SIDE OF CONFINING LAYER FOR LEAKAGE B.C. VISOAR - VISCOSITY IN OUTER AQUIFER REGION FOR A.I.F.B.C. VISTFO - VISCOSITY VS. TEMPERATURE DATA AT MINIMUM SOLUTE CONCENTRATION VISTER - VISCOSITY VS. TEMPERATURE DATA AT MAXIMUM SOLUTE CONCENTRATION VPA. - VARIABLY PARTITIONED ARRAY - REFERENCE VISCOSITY VREF **VSTLOG - LOGARITHM OF VISCOSITY** - INTERSTITIAL VELOCITY IN THE X-DIRECTION VXX VYY - INTERSTITIAL VELOCITY IN THE Y-DIRECTION VZZ - INTERSTITIAL VELOCITY IN THE Z-DIRECTION ₩0 - MINIMUM MASS FRACTION FOR SCALING 41 - MAXIMUM MASS FRACTION FOR SCALING 1800 - WELL-BORE OUTER DIANETER **HCAIF** - CUMULATIVE INFLOW AT AQUIFER INFLUENCE FUNCTION B.C. NCF - WELL COMPLETION FACTOR FOR A GIVEN LAYER WCLBL1 - LABEL DESCRIBING WELL CALCULATION TYPE WCLBL2 - LABEL DESCRIBING WELL-FLOW ALLOCATION NETHOD WFICUM - CUMULATIVE FLUID INJECTED BY A WELL WFPCUN - CUMULATIVE FLUID PRODUCED BY A WELL WHICUM - CUMULATIVE HEAT INJECTED BY A WELL HHPCUH - CUMULATIVE HEAT PRODUCED BY A HELL LI. - WELL INDEX WIDLBL - WELL IDENTIFICATION LABEL FOR TEMPORAL PLOT WOMETH - INDEX OF WELL-FLOW CALCULATION METHOD WRANGL - WELL-RISER ANGLE WITH THE GRAVITATIONAL VECTOR WRCALC - TRUE IF WELL-RISER CALCULATIONS ARE TO BE PERFORMED WRID - WELL-RISER PIPE INSIDE DIAMETER WRIDT - WELL-RISER PIPE INSIDE DIANETER FOR A GIVEN WELL WRISL - WELL-RISER PIPE LENGTH WRRUF - WELL-RISER PIPE ROUGHNESS PARAMETER WSICUM - CUMULATIVE SOLUTE INJECTED BY A WELL WSPCUM - CUNULATIVE SOLUTE PRODUCED BY A WELL - WEIGHT FACTOR FOR SPATIAL DISCRETIZATION UT. X - X-COORDINATE NODE LOCATIONS XC - X-COORDINATE LOCATION OF CALCULATED DATA FOR TEMPORAL PLOT XI - DIMENSIONLESS TIME PARAMETER FOR HEAT TRANSFER FROM WELL RISER - LABEL FOR X-AXIS FOR TEMPORAL PLOTS XLBL - X-COORDINATE LOCATION OF OBSERVED DATA FOR TEMPORAL PLOTS XÔ - SOLUTION VECTOR FROM TWO-LINE SOR SOLVER XX XXN - NEW SOLUTION VECTOR FROM TWO-LINE SOR SOLVER - Y-COORDINATE NODE LOCATIONS Y

- YC Y-COORDINATE LOCATIONS OF CALCULATED DATA FOR TEMPORAL PLOTS
- YLBL LABEL FOR Y-AXIS FOR TEMPORAL PLOTS
- YO Y-COORDINATE LOCATION OF OBSERVED DATA FOR TEMPORAL PLOTS
- YPOSUP TRUE IF THE Y-AXIS IS POSITIVE UPWARD FOR MAPS
- YY DEPENDENT VARIABLE VECTOR FOR WELL RISER INTEGRATION
- Z Z-COORDINATE NODE LOCATIONS
- ZCHARS CHARACTER ARRAY FOR ZEBRA-STRIPED CONTOUR MAPS
- ZEBRA TRUE IF ZONED CONTOUR MAPS ARE TO BE ZEBRA STRIPED WITH ALTERNATING SYMBOL AND BLANK ZONES
- ZELBC ELEVATION OF THE OUTER SURFACE OF A CONFINING LAYER FOR A LEAKAGE B.C.
- ZHCBC NODE LOCATIONS OUTSIDE THE REGION IN A NORMAL DIRECTION TO A HEAT CONDUCTION BOUNDARY FACE FOR HEAT CONDUCTION B.C.
- ZPINIT Z-COORDINATE LOCATION OF SPECIFIED INITIAL PRESSURE FOR HYDROSTATIC INITIAL CONDITIONS
- ZPOSUP TRUE IF THE Z-AXIS IS POSITIVE UPWARD FOR MAPS
- ZT Z-COORDINATE LOCATIONS OF TEMPERATURE PROFILE DATA FOR INITIAL CONDITIONS
- ZINC LOCATIONS ALONG THE OUTWARD NORMAL TO HEAT-CONDUCTION BOUNDARY FACES OF TEMPERATURE PROFILE DATA FOR INITIAL CONDITIONS
- ZWK LOCATION ALONG A WELL-RISER PIPE
- 2WKT Z-COORDINATE LOCATION OF A WELL DATUM

## 11.2 CROSS-REFERENCE LIST OF VARIABLES

The following cross-reference list (table 11.2) shows in which subprograms each variable appears. Actual line numbers within each subprogram are not provided; however, every FORTRAN compiler can provide a local crossreference map for a given subprogram. The first column contains the variable names and the second column lists the subprograms that employ each variable. The third column gives the variable type and the dimension, if an array. Some variables are scalars in some subprograms and arrays in others; however, the definitions are the same.

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
A	MAP2D	REAL*8 DIMENSION(*,*)	AAIF	APLYBC	REAL*8 DIMENSION(*)
	1	-		ASEMBL	REAL*8 DIMENSION(*)
AO	MAP2D	REAL*8		SUMCAL	REAL*8 DIMENSION(*)
A1	BSODE	REAL*8	ABOAR	HST3D	REAL*8
	INTERP	REAL*8		APLYBC	REAL*8
	MAP2D	REAL*8		DUMP	REAL*8
		\$ * -		ETOM1	REAL*8
A1HC	APLYBC	REAL*8 DIMENSION(*)		INIT2	REAL*8
	INIT2	REAL*8 DIMENSION(*)		READ1	REAL*8
				READ2	REAL*8
A2	BSODE	REAL*8		WRITE2	REAL*8
	INTERP	REAL*8			. · · · · ·
			ABPM	ETOM1	REAL*8 DIMENSION(*)
A2HC	APLYBC	REAL*8 DIMENSION(*)		INIT2	REAL*8 DIMENSION (*)
	INIT2	REAL*8 DIMENSION(*)		READ2	REAL*8 DIMENSION (*)
				WRITE2	REAL*8 DIMENSION(*)
A3HC	APLYBC	REAL*8 DIMENSION(*)			
	INIT2	REAL*8 DIMENSION(*)	ACHR	MAP2D	REAL*8
A4	D4DES	REAL*8 DIMENSION(*)	ALBC	APLYBC	REAL*8 DIMENSION(*)
	•			ASEMBL	REAL*8 DIMENSION(*)
AA1	APLYBC	REAL*8 DIMENSION(*)		SUMCAL	REAL*8 DIMENSION(*)
AA2	APLYBC	REAL*8 DIMENSION(*)	ALLOUT	ERROR2	LOGICAL*4
AA3	APLYBC	REAL*8 DIMENSION(*)	ALPHL	COEFF	REAL*8 DIMENSION(*)
,				ETOM1	REAL*8 DIMENSION(*)
AA4	APLYBC	REAL*8 DIMENSION(*)		READ2	REAL*8 DIMENSION(*)
				WRITE2	REAL*8 DIMENSION(*)

Table 11.2--Cross-reference list of variables

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
AHAX	HST3D	REAL*8 DIMENSION(3)	APLOT	HAP2D	REAL*8 DIMENSION (* ,*
	ETOM2	REAL*8 DIMENSION(3)			
	INIT3	REAL*8 DIMENSION(3)	APLOT 1	MAP2D	REAL*8
	MAP2D	REAL*8			
	READ3	REAL*8 DIMENSION(3)	APLOT2	MAP2D	REAL*8
	WRITE3	REAL*8 DIMENSION(3)			
	WRITES	REAL*8 DIMENSION(3)	APLOT3	HAP2D	REAL*8
AMAXN	MAP2D	REAL*8	APLOT4	MAP2D	REAL*8
AMIN	HST3D	REAL*8 DIMENSION(3)	APRNT	PRNTAR	REAL*8 DIMENSION(10)
	ETOM2	REAL*8 DIMENSION(3)		WRITE2	REAL*8 DIMENSION(*)
	INIT3	REAL*8 DIMENSION(3)		WRITE3	REAL*8 DIMENSION(*)
	MAP2D	REAL*8		WRITES	REAL*8 DIMENSION(*)
	READ3	REAL*8 DIMENSION(3)			
	WRITE3	REAL*8 DIMENSION(3)	ARGRID	HST3D	LOGICAL*4
	WRITE5	REAL*8 DIMENSION(3)		INIT2	LOGICAL*4
				READ2	LOGICAL*4
AMINN	HAP2D	REAL*8		WRITE2	LOGICAL*4
AMN	MAP2D	REAL*8	ARRAY	PRNTAR	REAL*8 DIMENSION(*)
AMX	HAP2D	REAL*8	ARWB	WELLSS	REAL*8
ANGOAR	HST3D	REAL*8	ARX	COEFF	REAL*8 DIMENSION(*)
	APLYBC	REAL*8		INIT2	REAL*8 DIMENSION(*)
	DUHP	REAL*8			
	ETOH1	REAL*8	ARXBC	INIT2	REAL*8 DIMENSION(*)
	INIT2	REAL*8			
	READ1	REAL*8	ARXFBC	INIT2	REAL*8 DIMENSION(*)
	READ2	REAL*8		INIT3	REAL*8 DIMENSION(*)
	WRITE2	REAL*8			

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
ARY	COEFF	REAL*8 DIMENSION(*)	BAIF	APLYBC	REAL*8 DIMENSION(*)
	INIT2	REAL*8 DIMENSION(*)		ASEMBL	REAL*8 DIMENSION(*)
	. •	• • • • •		SUMCAL	REAL*8 DIMENSION(*)
ARYBC	INIT2	REAL*8 DIMENSION(*)			
ARYFBC	INIT2	REAL*8 DIMENSION(*)	BBAIF	HST3D	REAL*8 DIMENSION(0:3)
	INIT3	REAL*8 DIMENSION(*)		APLYBC	REAL*8 DIMENSION (0:3)
				BLOCKDATA	
ARZ	COEFF	REAL*8 DIMENSION(*)		DUMP	REAL*8 DIMENSION(0:3)
	INIT2	REAL*8 DIMENSION(*)		READ1	REAL*8 DIMENSION(0:3)
				READ2	REAL*8 DIMENSION(0:3)
ARZBC	INIT2	REAL*8 DIMENSION(*)		WRITE2	REAL*8 DIMENSION(0:3)
ARZFBC	INIT2	REAL*8 DIMENSION(*)	BBLBC	APLYBC	REAL*8 DIMENSION(*)
	INIT3	REAL*8 DIMENSION(*)		ASEMBL	REAL*8 DIMENSION(*)
			1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 -	INIT2	REAL*8 DIMENSION(*)
AST	PLOT	CHARACTER*1		SUMCAL	REAL*8 DIMENSION(*)
	ZONPLT	CHARACTER*1		WRITE2	REAL*8 DIMENSION(*)
AUTOTS	HST3D	LOGICAL*4	BETA	APLYBC	REAL*8 DIMENSION(10)
	COEFF	LOGICAL*4			
	ERROR3	LOGICAL*4	BLANK	MAP2D	CHARACTER*1
	ETOM2	LOGICAL*4		PLOT	CHARACTER*1
	INIT3	LOGICAL*4		PRNTAR	CHARACTER*1
	READ3	LOGICAL*4		<b>WRITE3</b>	CHARACTER*11
	WRITE3	LOGICAL*4		ZONPLT	CHARACTER*1
BO	BSODE	REAL*8	BLANKL	PLOT	CHARACTER*51
B1	BSODE	REAL*8	BLBC	APLYBC	REAL*8 DIMENSION(*)
-	WFDYDZ	REAL*8		ASEMBL SUMCAL	REAL*8 DIMENSION(*) REAL*8 DIMENSION(*)
B2	WFDYDZ	REAL*8			• •

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable Lype	Variable name	Refer- encing programs	Variable type
BOAR	HST3D	REAL*8	BV1	HST3D	REAL*8
	APLYBC	REAL*8		DUMP	REAL*8
	DUMP	REAL*8		INIT2	REAL*8
	ETOH1	REAL*8		READ1	REAL*8
	INIT2	REAL*8		VISCOS	REAL*8
	READ1	REAL*8			
	READ2	REAL*8	С	APLYBC	REAL*8 DIMENSION(*)
	WRITE2	REAL*8		ASEMBL	REAL*8 DIMENSION(*)
<b>n</b> n			•	CALCC	REAL*8
BP	list3D	REAL*8		COEFF	REAL*8 DIMENSION(*)
	DUMP	REAL*8		CRSDSP	REAL*8 DIMENSION(*)
	ETOH1	REAL*8		INIT2	REAL*8 DIMENSION(*)
	INIT2	REAL*8		ITER	REAL*8 DIMENSION(*)
•	READ1	REAL*8		READ2	REAL*8 DIMENSION(*)
	READ2	REAL*8		SUHCAL	REAL*8 DIMENSION(*)
	WRITE2	REAL*8		VISCOS	REAL*8
1) <b>M</b>	10000			WBBAL	REAL*8 DIMENSION(*)
BT	HST3D	REAL*8		WELLSS	REAL*8 DIMENSION(*)
	DUMP	REAL*8		WRITE2	REAL*8 DIMENSION(*)
	ETOH1	REAL*8		WRITE5	REAL*8 DIMENSION(*)
	INIT2	REAL#8		ZONPLT	REAL*8
	READ1	REAL*8			
	READ2	REAL*8	C00	HST3D	REAL*8
	WRITE2	REAL*8		WBBAL	REAL*8
BV	UCTNER	DRAT-LO		WELLSS	REAL*8
DV	VSINIT	REAL*8		WELRIS	REAL*8
BVO	UCTON	DEALS		WFDYDZ	REAL*8
DAA	HST3D	REAL*8			
	DUMP INIT2	REAL*8	C1	ZONPLT	REAL*8
		REAL*8			<b>-</b> ,
	READ1	REAL*8		•	
	VISCOS	REAL*8			

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
C11	HST3D	REAL#8	C31	HST3D	REAL*8
	ASEMBL	REAL*8		ASEMBL	REAL*8
	CALCC	REAL*8		CALCC	REAL*8
	WFDYDZ	REAL*8	,	•	en e
			C32	HST3D	REAL*8
C12	HST3D	REAL*8	· · · ·	ASEMBL	REAL*8
	ASEMBL	REAL*8		CALCC	REAL*8
	CALCC	REAL*8			n e an
	WFDYDZ	REAL*8	C33	HST3D	REAL*8
•				ASEMBL	REAL#8
C13	HST3D	REAL*8		CALCC	REAL*8
	ASEMBL	REAL*8		1	· · ·
	CALCC	REAL*8	C34	HST3D	REAL*8
C2	ZONPLT	REAL*8	2	ASEMBL	REAL#8
				CALCC	REAL*8
C21	HST3D	REAL*8			
	ASEMBL	REAL*8	C35	HST3D	REAL*8
	CALCC	REAL*8		ASEMBL	REAL*8
	WFDYDZ	REAL*8		CALCC	REAL*8
· ·	•				•
C22	HST3D	REAL*8	CAIF	APLYBC	REAL*8 DIMENSION(*
	" ASEMBL	REAL*8		ASEMBL	REAL*8 DIMENSION(*)
	CALCC	REAL*8		INIT3	REAL*8 DIMENSION (*)
:	WFDYDZ	REAL*8		SUMCAL	REAL*8 DIMENSION(*)
C23	HST3D	REAL*8	CC24	ASEMBL	REAL*8 DIMENSION (*
· · ·	ASEMBL	REAL*8			
	CALCC	REAL*8	CC34	ASEMBL	REAL*8 DIMENSION(*
C24	HST3D	REAL*8	CC35	ASEMBL	REAL*8 DIMENSION (*
	ASEMBL	REAL*8			
	CALCC	REAL*8			•

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable statype	Variable name	Refer- encing programs	Variable type
CCLBL	PLOTOC	CHARACTER*50	CIBC	APLYBC	CHARACTER*9
				ASEMBL	CHARACTER*9
CCW	PLOTOC	REAL*8 DIMENSION(*)		ERROR2	CHARACTER*9
				ERROR3	CHARACTER*9
CFLX	APLYBC	REAL*8 DIMENSION(*)		ETOH1	CHARACTER*9
	INIT3	REAL*8 DIMENSION(*)		INIT2	CHARACTER*9
	SUMCAL	REAL*8 DIMENSION(*)		INIT3	CHARACTER*9
	WRITE3	REAL*8 DIMENSION(*)		SBCFLO	CHARACTER*9
	WRITES	REAL*8 DIMENSION(*)		SUMCAL	CHARACTER*9
				WBCFLO	CHARACTER*9
CHAPRT	WRITE5	CHARACTER*12 DIMENSION(10)		WRITE2	CHARACTER*9
				WRITE3	CHARACTER*9
CHARC	READ1	CHARACTER*1		WRITE5	CHARACTER*9
CHARS	HAP2D	CHARACTER*1 DIMENSION(0:31)	CIBC1	SOR2L	CHARACTER*9
CHK1	WRITE5	CHARACTER*2	CIBC2	SOR2L	CHARACTER*9
CHKPTD	HST3D	LOGICAL#4	CICALL	IREWI	CHARACTER*1
	CLOSE	LOGICAL*4			,
	DUMP	LOGICAL*4	CLBC	APLYBC	REAL*8 DIMENSION(*)
	READ3	LOGICAL#4		ASEHBL	REAL*8 DIMENSION(*)
				INIT3	REAL*8 DIMENSION(*)
Chu3	WRITE5	CHARACTER*12		SUMCAL	REAL*8 DIMENSION(*)
				WRITE3	REAL*8 DIMENSION(*)
CHU4	WRITES	CHARACTER*8			
	•		CLINE	HAP2D	REAL*8
CHU5	WRITES	CHARACTER*8			
-			CMAX	PLOTOC	REAL*8
CHU6	WRITE5	CHARACTER*8			
			CHIN	PLOTOC	REAL*8
CI	D4DES	INTEGER*4 DIMENSION(6,*)			
	ORDER	INTEGER*4 DIMENSION(6,*)	CHX	ASEMBL	REAL*8

Table 11.2--Cross-reference list of variables--Continued

Variable nam <del>e</del>	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
СМУ	ASEMBL	REAL*8	CNVDI	HST3D	REAL*8
		•		DUMP	REAL*8
CMZ	ASEMBL	REAL*8		INIT1	REAL*8
				PLOTOC	REAL*8
CN3	PRNTAR	CHARACTER*2		READ1	REAL*8
				WRITE2	REAL*8
CNP	CALCC	REAL*8		WRITE3	REAL*8
	READ3	REAL*8 DIMENSION(*)		WRITE4	REAL*8
				WRITE5	REAL*8
CNV	ETOM1	REAL*8			• •
	PRNTAR	REAL*8	CNVE	ETOM2	REAL*8
	WRITE2	REAL*8			
	WRITE3	REAL*8	CNVEPI	HST3D	REAL*8
	WRITE5	REAL*8		DUMP	REAL*8
				INIT1	REAL*8
CNVD	ETOMÍ	REAL*8		READ1	REAL*8
	ETOM2	REAL*8		WRITE3	REAL*8
	READ3	REAL*8			
			CNVFF	ETOM2	REAL*8
CNVDFI	HST3D	REAL*8		READ3	REAL*8
	DUMP	REAL*8			
	INIT1.	REAL*8	CNVGZ	READ3	REAL*8
	PLOTOC	REAL*8			
	READI	REAL*8	CNVHC	ETOM1	REAL*8
	WRITE2	REAL*8		ETOM2	REAL*8
	WRITE3	REAL+8			
	WRITE4	REAL*8		1	
	WRITE5	REAL <sup>+8</sup>			

Table 11.2--Cross-reference list of variables--Continued

/ariable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
CNVIICI	HST3D	REAL*8	CNVIITC	ETOHI	REAL*8
	DUMP	REAL*8		ETOH2	REAL*8
	INITI	REAL*8		. DIGIL	
	PLOTOC	REAL*8	CNVHTI	WRITE2	REAL*8
	READ1	REAL*8			
	WRITE2	REAL*8	CNVL	ETOM1	REAL*8
	WRITE3	REAL*8		READ3	REAL*8
	WRITE4	REAL*8			
	WRITES	REAL*8	CNVL2	ETOH1	REAL*8
				ETOM2	REAL*8
CNVHF	ETOM2	REAL*8		READ3	REAL*8
	READ3	REAL*8			
			CNVL21	HST3D	REAL*8
CNVIIFI	list3D	REAL*8	,	DUMP	REAL*8
	DUMP	REAL*8		INIT1	REAL*8
	INITI	REAL*8		PLOTOC	REAL*8
	PLOTOC	REAL*8		READ1	REAL*8
	READ1	REAL*8		WRITE2	REAL*8
	WRITE2	REAL*8		WRITE3	REAL*8
	WRITE3	REAL*8		WRITE4	REAL*8
	WRITE4	REAL*8		WRITE5	REAL*8
	WRITE5	REAL*8	<b>ANR 17 A</b>		
			CNVL3	ETOM1	REAL*8
CNVHI	HST3D	REAL*8		ETOM2	REAL*8
	DUMP	REAL*8	01011 0 1	110000	
	IN <b>IT1</b>	REAL*8	CNVL31	HST3D	REAL*8
	PLOTOC	REAL*8		DUMP	REAL*8
	READ1	REAL*8		INIT1	REAL*8
	WRITE2	REAL*8		PLOTOC	REAL*8
	WRITE3	REAL*8		READ1	REAL*8
	WRITE4	REAL*8		WRITE2	REAL*8
	WRITE5	REAL*8		WRITE3	REAL*8
				WRITE4	REAL*8
*				WRITE5	REAL*8
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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
CNVLI	HST3D	REAL*8	CNVP	ETOM1	REAL*8
	DUMP	REAL#8		ETOM2	REAL*8
	INITI	REAL*8		READ3	REAL*8
	PLOTOC	REAL*8			
	READ1	REAL*8	CNVPI	HST3D	REAL*8
	WRITE2	REAL*8		DUMP	REAL*8
	WRITE3	REAL*8		INITI	REAL*8
	WRITE4	REAL*8		PLOTOC	REAL*8
	WRITES	REAL*8		READ1	REAL*8
				WRITE2	REAL*8
CNVMFI	listod	REAL*8		WRITE3	REAL*8
	DUMP	REAL*8		WRITE4	REAL*8
	INIT1	REAL*8		WRITE5	REAL*8
	PLOTOC	REAL*8			1. A.
	READ1	REAL*8	CNVSF	ETOM2	REAL*8
	WRITE2	REAL*8		READ3	REAL*8
	WRITE3	REAL*8			
	WRITE4	REAL*8	CNVT1	ETOM1	REAL*8
	WRITE5	REAL*8		ETOM2	REAL*8
				READ3	REAL*8
CNVMI	HST3D	REAL*8			
<b>-</b>	DUMP .	REAL*8	CNVT1I	HST3D	REAL*8
	INITI	REAL*8		DUMP	REAL*8
	PLOTOC	REAL*8		INITI	REAL*8
	READ1	REAL*8		PLOTOC	REAL*8
1	WRITE2	REAL*8	, ·	READ1	REAL*8
	WRITE3	REAL*8		WRITE2	REAL*8
	WRITE4	REAL*8		WRITE3	REAL*8
	WRITE5	REAL*8		WRITE4	REAL*8
	•••••		. •	WRITE5	REAL*8

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
CNVT2	ETOM1	REAL*8	CNVTHI	HST3D	REAL*8
	ETOM2	REAL*8		DUMP	REAL*8
	READ3	REAL*8		INIT1	REAL*8
				PLOTOC	REAL*8
CNVT2I	HST3D	REAL*8	•	READ1	REAL*8
	DUHP	REAL*8		WRITE2	REAL*8
	INIT1	REAL*8		WRITE3	REAL*8
	PLOTOC	REAL*8		WRITE4	REAL*8
	READ1	REAL*8		WRITE5	REAL*8
	WRITE2	REAL*8			
	WRITE3	REAL*8	CNVUUI	WRITE4	REAL*8
	WRITE4	REAL*8			
	WRITES	REAL*8	CNVVF	етон1	REAL*8
				READ3	REAL*8
CNVTCI	HST3D	REAL*8			
	DUMP	REAL*8	CNVVLI	HST3D	REAL*8
	init1	REAL*8		DUMP	REAL*8
	PLOTOC	REAL*8		INIT1	REAL*8
	READ1	REAL*8		PLOTOC	REAL*8
	WRITE2	REAL*8		READ1	REAL*8
	WRITE3	REAL*8		WRITE2	REAL*8
	WRITE4	REAL*8		WRITE3	REAL*8
	WRITES	REAL*8		WRITE4	REAL*8
				WRITES	REAL*8
CNVTHC	ETOM1	REAL*8			
	etom2	REAL*8	CNVVSI	HST3D	REAL*8
				DUMP	REAL*8
CNVTH	ETOM1	REAL*8		INITI	REAL*8
	ETOM2	REAL*8		PLOTOC	REAL*8
	READ1	REAL*8		READ1	REAL*8
				WRITE?	REAL*8
				WRITE3	REAL*8
				WRITE4	REAL*8
				WRITES	REAL*8

Table 11.2--Cross-reference list of variables--Continued

Variable nam <del>e</del>	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
COLS	MAP2D	REAL*8	CPF	HST3D	REAL*8
				APLYBC	REAL*8
COMOPT	ITER	LOGICAL*4		ASEMBL	REAL*8
	L2SOR	LOGICAL*4		CALCC	REAL*8
				COEFF	REAL*8
CONLBL	WRITES	CHARACTER*20		CRSDSP	REAL*8
	· · ·			DUMP	REAL*8
CONVC	HST3D	LOGICAL*4		ETOM1	REAL*8
	ITER	LOGICAL*4		INIT2	REAL*8
		Ŷ		READ1	REAL*8
CONVP	HST3D	LOGICAL*4		READ2	REAL*8
	ITER	LOGICAL*4		SUMCAL	REAL*8
				WELLSS	REAL*8
CONVRG	BSODE	LOGICAL*4		WELRIS	REAL*8
				WFDYDZ	REAL*8
CONVT	HST3D	LOGICAL*4		WRITE2	REAL*8
4 4 *	ITER	LOGICAL*4		WRITE5	REAL*8
COW	PLOTOC	REAL*8 DIMENSION(*)	СРХ	ASEMBL	REAL*8
CPAR	IREWI	CHARACTER*9	СРУ	ASEMBL	REAL*8
CPARI	IREWI	CHARACTER*6	CPZ .	ASEMBL	REAL*8

Table 11.2--Cross-reference list of variables--Continued

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Variable	Refer- encing	V	ariable	Variable	Refer- encing	Variable
name	programs		type	name	programs	type
CSBC	ASEMBL	REAL*8	DIMENSION(*)	CYLIND	HST3D	LOGICAL*4
	INIT3	REAL*8	DIMENSION(*)		ASEHBL	LOGICAL*4
	SUMCAL	REAL*8	DIMENSION (*)		CLOSE	LOGICAL*4
	WRITE3	REAL*8	DIMENSION(*)		COEFF	LOGICAL*4
					DUHP	LOGICAL*4
CVIS	HST3D	REAL*8	DIMENSION(10)		ERROR2	LOGICAL*4
	DUMP		DIMENSION(10)		ERROR3	LOGICAL*4
	ETOM1	REAL*8	DIMENSION(10)		ETOM1	LOGICAL*4
	INIT2	REAL*8	DIMENSION(10)		INITI	LOGICAL*4
	READ1	REAL*8	DIMENSION(10)		INIT2	LOGICAL*4
	READ2	REAL*8	DIMENSION(10)		INIT3	LOGICAL*4
	VISCOS	REAL*8	DIMENSION(10)		IREWI	Logical*4
	WRITE2	REAL*8	DIMENSION(10)		ITER	LOGICAL*4
					L2SOR	LOGICAL*4
CWKT	ASEMBL	REAL*8	DIMENSION(*)		PLOTOC	LOGICAL*4
	READ3	REAL*8	DIMENSION(*)		READ1	LOGICAL*4
	WBBAL	REAL*8	DIMENSION (*)		READ2	LOGICAL*4
	<b>WELLSS</b>	REAL*8	dimension(*)		READ3	LOGICAL*4
	WRITE3	REAL*8	DIMENSION(*)		REWI	LOGICAL*4
	WRITE5	REAL*8	dimension(*)		REWI3	Logical*4
					SUMCAL	LOGICAL*4
					WBBAL	LOGICAL*4
					WELLSS	LOGICAL*4
					WRITE1	LOGICAL*4
					WRITE2	LOGICAL*4
					WRITE3	LOGICAL*4
					WRITE4	LOGICAL*4
					<b>WRITES</b>	LOGICAL*4
					ZONPLT	LOGICAL*4

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Variable nam <del>e</del>	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
D	SOR2L	REAL*8 DIMENSION(*)	DCTAS	HST3D	REAL*8
		ι ·		COEFF	REAL*8
DAMWRC	HST3D	REAL*8		ERROR3	REAL*8
	DUMP	REAL*8		ETOM2	REAL*8
	INIT2	REAL*8		INIT3	REAL*8
	READ1	REAL*8	•	READ3	REAL*8
	READ2	REAL*8		WRITE3	REAL*8
	WELLSS	REAL*8			
	WRITE2	REAL*8	DDNMAX	HST3D	REAL*8
		·		SUMCAL	REAL*8
DASHES	WRITE5	CHARACTER*50		WRITE5	REAL*8
DBKD	INIT2	REAL*8 DIMENSION(*)	DDV	SBCFLO	REAL*8 DIMENSION(*)
· · · · ·	READ2	REAL*8 DIMENSION(*)		WBCFLO	REAL*8 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(*)			
			DECLAM	HST3D	REAL*8
DC	ASEMBL	REAL*8 DIMENSION(*)		ASEMBL	REAL*8
	CALCC	REAL*8		CALCC	REAL*8
	CRSDSP	REAL*8 DIMENSION(*)		COEFF	REAL*8
	ITER	REAL*8 DIMENSION(*)		DUMP	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)		ETOM1	REAL*8
				INIT2	REAL*8
DCMAX .	HST3D	REAL*8		INIT3	REAL*8
	COEFF	REAL*8		READ1	REAL*8
	ITER	REAL*8		READ2	REAL*8
	SUMCAL	REAL*8	•	READ3	REAL*8
۰.	WRITE5	REAL*8		SUMCAL	REAL*8
				VISCOS	REAL*8
				WBBAL	REAL*8
	1 - 1			WELLSS	REAL*8
				WRITE2	REAL*8
				WRITE3	REAL*8
				WRITE5	REAL*8

Table 11.2--Cross-reference list of variables--Continued

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	encing programs	Variable <b>type</b>	Variable name	Refer- encing programs	Variable type
DEHIR	HST3D	REAL*8	DEN	APLYBC	REAL*8 DIMENSION(*)
	SUHCAL	REAL*8		ASEHBL	REAL*8 DIMENSION(*)
	WRITES	REAL*8		COEFF	REAL*8 DIMENSION(*)
				INIT2	REAL*8 DIMENSION(*)
DELA	HAP2D	REAL*8		SUHCAL	REAL*8 DIMENSION(*)
				WBBAL	REAL*8 DIMENSION(*)
DELAN	MAP2D	REAL*8		WELLSS	REAL*8 DIMENSION(*)
				WRITE2	REAL*8 DIMENSION(*)
DELAP	MAP2D	REAL*8		WRITE5	REAL*8 DIMENSION(*)
DELTIM	HST3D	REAL*8	DENO	HST3D	REAL*8
	APLYBC	REAL*8		CALCC	REAL*8
	CALCC	REAL*8		DUMP	REAL*8
	COEFF	REAL*8		INIT2	REAL*8
	DUMP	REAL*8		ITER	REAL*8
	ETOM2	REAL*8		READ1	REAL*8
	INIT2	REAL*8		SUHCAL	REAL*8
	INIT3	REAL*8		WBBAL	REAL*8
	READ1	REAL*8		WELLSS	REAL*8
	READ3	REAL*8		WFDYDZ	REAL*8
	SUHCAL	REAL*8			
	WELLSS	REAL*8	DENC	HST3D	REAL*8
	WRITE3	REAL*8		CALCC	REAL*8
	WRITE4	REAL*8		DUMP	REAL*8
	WRITE5	REAL*8		INIT2	REAL*8
				ITER	REAL*8
DELX	INIT2	REAL*8		READ1	REAL*8
				SUMCAL	REAL*8
DELY	INIT2	REAL*8		WBBAL	REAL*8
	PLOT	REAL*8		WELLSS	REAL*8
				WFDYDZ -	REAL*8
DELZ	INIT2	REAL*8	DENCIIC	ITER	REAL*8

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
DENCHT	ITER	REAL*8	DENMAX	SUMCAL	REAL*8
DENFO	HST3D	REAL*8	DENN	CALCC	REAL*8
	DUMP	REAL*8			
ETOM1 INIT2	ETOM1	REAL*8	DENNP	CALCC	REAL*8
	INIT2	REAL*8			
	READ1	REAL*8	DENOAR	APLYBC	REAL*8 DIMENSION(*)
	READ2	REAL*8		ASEMBL	REAL*8 DIMENSION(*)
	WRITE2	REAL*8		ETOM1	REAL*8
				INIT3	REAL*8 DIMENSION(*)
DENF1 I	HST3D	REAL*8		SUMCAL	REAL*8 DIMENSION(*)
	DUMP	REAL*8			
	ETOM1	REAL*8	DENP	HST3D	REAL*8
. r	INIT2	REAL*8		CALCC	REAL*8
	READ1	REAL*8		DUMP	REAL*8
	READ2	REAL*8		INIT2	REAL*8
	WRITE2	REAL*8		ITER	REAL*8
				READ1	REAL#8
DENFBC	APLYBC	REAL*8 DIMENSION(*)		SUMCAL	REAL*8
	INIT3	REAL*8 DIMENSION(*)		WBBAL	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)		WELLSS	REAL#8
	WRITE3	REAL*8 DIMENSION(*)		WFDYDZ	REAL*8
	WRITES	REAL*8 DIMENSION(*)			
		· · · ·	DENT	HST3D	REAL*8
DENGL	WELLSS	REAL*8		CALCC	REAL*8
				DUMP	REAL*8
DENLBC	APLYBC	REAL*8 DIMENSION(*)		INIT2	REAL*8
	ASEMBL	REAL*8 DIMENSION(*)		ITER	REAL*8
	INIT3	REAL*8 DIMENSION(*)		READ1	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)		SUMCAL	REAL*8
	WRITE3	REAL*8 DIMENSION (*)		WBBAL	REAL*8
				WELLSS	REAL*8
				WFDYDZ	REAL*8

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
DENW	WELLSS	REAL*8 DIMENSION(*)	DOTS	CLOSE	CHARACTER*50
				REWI	CHARACTER*50
DENWKT	ASEMBL	REAL <sup>2</sup> 8		REWI3	CHARACTER*50
	WBBAL	REAL*8	•	WRITE2	CHARACTER*50
	WELLSS	REAL*8		WRITE3	CILARACTER*50
				WRITE5	CHARACTER*50
DENWRK	HST3D	REAL*8			
	WELRIS	REAL*8	DP	ASEMBL	REAL*8 DIMENSION(*)
	WFDYDZ	REAL*8		CALCC	REAL*8
				ITER	REAL*8 DIMENSION(*)
DET	WFDYDZ	REAL*8		SUNCAL	REAL*8 DIMENSION(*)
				WBBAL	REAL*8 DIMENSION(*)
DFIR	HST3D	REAL*8		WRITE5	REAL*8 DIMENSION(*)
	SUHCAL	REAL*8			
	WRITE5	REAL*8	DPMAX	hst3d	REAL*8
DM	HST3D	REAL#8		COEFF	REAL*8
Uri -	ASEMBL	REAL*8		ITER	REAL*8
	COEFF	REAL*8		SUHCAL	REAL*8
	DUMP	REAL*8	i.	WRITE5	REAL*8
	ETOM1	REAL*8			
	INIT2	REAL*8	DPTAS	HST3D	REAL*8
	INIT3	REAL*8		COEFF	REAL*8
	READ1	REAL*8		ERROR3	REAL*8
	READ2	REAL*8		etom2	REAL*8
	READ2 READ3	REAL=0 REAL*8		INIT3	REAL*8
	SUMCAL	REAL*8		READ3	REAL*8
	VISCOS	REAL*8		WRITE3	REAL*8
	WBBAL	REAL*8			
	WELLSS	REAL*8	DPUDT	APLYBC	REAL*8
	WRITE2	REAL*8			
	WRITE3	REAL*8	DPWKT	ASEMBL	REAL*8 DIMENSION(*)
	WRITE5	REAL*8		i,ter	REAL*8 DIMENSION(*)
				WBBAL '	REAL*8 DIMENSION(*)
<b>`'TEST</b>	WELLSS	REAL*8	(	WRITE5	REAL*8 DIMENSION(*)
			(		

Table 11.2Cross-reference	list of variablesContinued
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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
DQFDP	ASEMBL	REAL*8	DQWLYR	ASEMBL	REAL*8
DQHBC	ASEMBL	REAL*8	DSIR	HST3D	REAL*8
	SUMCAL	- REAL*8		SUMCAL	REAL*8
				WRITES	REAL*8
DQHCDT	APLYBC	REAL*8 DIMENSION(*)			
	ASEMBL	REAL*8 DIMENSION(*)	DSXXM	ASEMBL	REAL*8
	ITER	REAL*8 DIMENSION(*)			
	SUMCAL	REAL*8 DIMENSION(*)	DSXXP	ASEMBL	REAL*8
DQHDP	ASEMBL	REAL*8	DSYYM	ASEMBL	REAL*8
DQHDT	ASEMBL	REAL*8	DSYYP	ASEMBL	REAL*8
DQHWDP	ASEMBL	REAL*8	DSZZM	ASEMBL	REAL*8
DQHWDT	ASEMBL	REAL*8	DSZZP	ASEMBL	REAL*8
DQSBC	ASEMBL	REAL*8	DT	ASEMBL	REAL*8 DIMENSION(*)
•	SUMCAL	REAL*8	. '	CALCC	REAL*8
				CRSDSP	REAL*8 DIMENSION(*)
DQSDC	ASEMBL	REAL*8		ITER	REAL*8 DIMENSION(*)
	•	·		SUMCAL	REAL*8 DIMENSION(*)
DQSDP	ASEMBL	REAL*8		· · .	
•.			DTADZW	HST3D	REAL*8
DQSWDC	ASEMBL	REAL*8		WELRIS	REAL*8
-				WFDYDZ	REAL*8
DQSWDP	ASEMBL	REAL*8			
		• •	DTHAWR	ETOM1	REAL*8 DIMENSION(*)
DQWDP	ASEMBL	REAL*8	· .	READ2	REAL*8 DIMENSION(*)
				WELRIS	REAL*8
DQWDPL	ASEMBL	REAL*8 DIMENSION(*)		WRITE2	REAL*8 DIMENSION(*)
	ITER	REAL*8 DIMENSION(*)	<u> </u>	•	
	WBBAL	REAL*8 DIMENSION(*)	DTHHC	APLYBC	REAL*8 DIMENSION(*)
	WELLSS	REAL*8 DIMENSION(*)		' INIT2	REAL*8 DIMENSION(*)

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing progr <b>ams</b>	Variable type	Variable name	Refer- encing programs	Variable type
DTIMMN	HST3D	REAL*8	DX	ZONPLT	REAL*8
	COEFF	REAL*8			
	ERROR3	REAL*8	DXMIN	PLOT	REAL*8
	ETOM2	REAL*8			
	INIT3	REAL*8	DXPRNT	PLOT	REAL*8
	READ3	REAL*8			
	WRITE3	REAL*8	DY	ZONPLT	REAL*8
DTIMOX	HST3D	REAL*8	DYY	BSODE	REAL*8 DIMENSION(2)
	COEFF	REAL*8		WELRIS	REAL*8 DIMENSION(2)
	ERROR3	REAL*8		WFDYDZ	REAL*8 DIMENSION(2)
	eton2	REAL*8		•	
	INIT3	REAL*8	DYYN	BSODE	REAL*8 DIMENSION(2)
	READ3	REAL*8			
	WRITE3	REAL*8	DZ	ASEMBL	REAL*8
				BSODE	REAL*8
DTHAX	HST3D	REAL*8		CALCC	REAL*8
	COEFF	REAL*8	•		
	ITER	REAL*8	DZCHNG	BSODE	REAL*8
	SUMCAL	REAL*8			
	<b>WRITES</b>	REAL*8	DZHIN	HST3D	REAL*8
				BSODE	REAL*8
DTTAS	HST3D	REAL*8		DUMP	REAL*8
	COEFF	REAL*8		eton1	REAL*8
	ERROR3	REAL*8		INIT2	REAL*8
	ETOM2	REAL*8		READ1	REAL*8
	INIT3	REAL*8		READ2	REAL*8
	READ3	REAL*8		WELRIS	REAL*8
	WRITE3	REAL*8			
			DZW	WELRIS	REAL*8

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Variable	Refer- encing	Variable	Variable	Refer- encing	Variable
Dauc	programs	type	name	programs	type
EEUNIT	HST3D	LOGICAL*4	EHO	HST3D	REAL*8
	ASEMBL	LOGICAL*4		APLYBC	REAL*8
	CLOSE	LOGICAL*4		ASEMBL	REAL*8
	COEFF	LOGICAL*4		DUMP	REAL*8
	DUMP	LOGICAL*4		INIT2	REAL*8
	ERROR2	LOGICAL*4		READ1	REAL*8
	ETOM1	LOGICAL*4		SUMCAL	REAL*8
	INIT1	LOGICAL*4	1	WBBAL	REAL*8
	INIT2	LOGICAL#4		WELLSS	REAL*8
	INIT3	LOGICAL*4		WRITE2	REAL*8
	IREWI	LOGICAL*4		WRITE5	REAL*8
	PLOTOC	LOGICAL*4			. •
	READ1	LOGICAL*4	EHOO	HST3D	REAL*8
	READ2	LOGICAL*4	• •	WBBAL	REAL*8
	READ3	LOGICAL*4		WELLSS	REAL*8
	REWI	LOGICAL*4		WELRIS	REAL*8
	REWI3	LOGICAL*4			
	SUMCAL	LOGICAL*4	11 A.		· · · ·
	WELLSS	LOGICAL*4	EHDT	HST3D	REAL*8 DIMENSION(14,10
	WRITE1	LOGICAL*4		BLOCKDATA	REAL*8 DIMENSION(14,10
	WRITE2	LOGICAL*4		DUMP	REAL*8 DIMENSION(14,10
	WRITE3	LOGICAL*4		INIT2	REAL*8 DIMENSION(14,10
	WRITE4	LOGICAL*4		READ1	REAL*8 DIMENSION(14,10
	WRITES	LOGICAL*4		TOFEP	REAL*8 DIMENSION(14,10
EH and a	APLYBC	REAL*8 DIMENSION(*)	EHI	TOFEP	REAL*8
	ASEMBL	REAL*8 DIMENSION(*)			
	COEFF	REAL*8 DIMENSION(*)	EHIR	HST3D	REAL*8
	INIT2	REAL*8 DIMENSION(*)		SUMCAL	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)		WRITES	REAL*8
	TOFEP	REAL*8			
	WBBAL	REAL*8 DIMENSION(*)		_	
	WELLSS	REAL*8 DIMENSION(*)			
	WRITE2	REAL*8 DIMENSION(*)			
		DOATHO DIMOTON (+)			

REAL\*8 DIMENSION(\*)

WRITE5

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
EHIRO	list3D	REAL*8	EHWKT	ASEMBL	REAL*8 DIMENSION(*)
	APLYBC	REAL*8		WBBAL	REAL*8 DIMENSION(*)
	DUMP	REAL*8		WELLSS	REAL*8 DIMENSION(*)
	INIT2	REAL*8		WRITE5	REAL*8 DIMENSION(*)
	READ1	REAL*8			
	SUHCAL	REAL*8	EIIWSUR	WBBAL	REAL*8 DIMENSION(*)
	WRITE2	REAL*8		WELLSS	REAL*8 DIMENSION(*)
	WRITES	REAL*8		WRITE5	REAL*8 DIMENSION(*)
EHIRN	SUMCAL	REAL*8	ELBL	PLOT	CHARACTER* 1
EIIMX	ASEMBL	REAL*8	ELO	TOFEP	REAL*8
EHMY	ASEMBL	REAL*8	ENTH	TOFEP	REAL*8
EIIMZ	ASEMBL	REAL*8	EOD	HST3D	REAL*8
				WELLSS	REAL*8
EHPX	ASEMBL	REAL*8		WELRIS	REAL*8
				WFDYDZ	REAL*8
EIIPY	ASEMBL	REAL*8			
			EODF	WELLSS	REAL*8
EHPZ	ASEMBL	REAL*8		WFDYDZ	REAL*8
EHST	HST3D	REAL*8 DIMENSION(32)	EPS	TOFEP	REAL*8
	BLOCKDATA				
	DUMP	REAL*8 DIMENSION(32)	EPSFAC	SOR2L	REAL*8
	INIT2	REAL*8 DIMENSION(32)			
	READ1	REAL*8 DIMENSION(32)	EPSONG	HST3D	REAL*8
	TOFEP	REAL*8 DIMENSION(32)		DUMP	REAL*8
				INIT2	REAL*8
Elivend	HST3D	REAL*8		READ1	REAL*8
	WBBAL	REAL*8		READ2	REAL*8
	WELLSS	REAL*8		SOR2L	REAL*8
	WELRIS	REAL*8			

Table 11.2--Cross-reference list of variables--Continued

EPSSOR	encing programs HST3D DUMP INIT2 READ1 READ2	Variable type REAL*8 REAL*8 REAL*8 REAL*8	Variable name ERREXI	encing programs HST3D	Variable type
EPSSOR	HST3D DUMP INIT2 READ1	REAL*8 REAL*8			·····
	DUMP INIT2 READ1	REAL*8	ERREXI	HST3D	
	INIT2 READ1			_	LOGICAL*4
· · · · · ·	READ1	RRAT*8		CLOSE	LOGICAL*4
· · · · · ·		STATUS V		INIT2	LOGICAL*4
	READ2	REAL#8	•	WRITE2	LOGICAL*4
×		REAL*8	EVEN	SOR2L	LOGICAL*4
	SOR2L	REAL*8	EVEN	SUKZL	INGICAL~4
	WRITE2	REAL*8	EVMAX	SOR2L	REAL*8
	HST3D	REAL*8	EVMIN	SOR2L	REAL*8
	BSODE	REAL*8	EAUTU	JUNZL	NGAL O
	DUMP	REAL*8	EX	PLOT	CHARACTER*1
	INIT2	REAL*8	<b>F1</b>		Guuriostin's
	READ1	REAL*8	EXTRAP	BSODE	REAL*8 DIMENSION(2,11)
	READ2	REAL*8	MUSIM	00000	THE O PERMOTON(L)
	WELRIS	REAL*8	F10	ZONPLT	CHARACTER*20
ERREX	ERROR 1	LOGICAL*4	F12	ZONPLT	CHARACTER*20
	ERROR2	LOGICAL*4	F 1 Z	LUNELI	CIPARCIES"20
	ERROR3	LOGICAL*4	FIAIF	HST3D	REAL*8
	INTERP	LOGICAL*4	E MIE	APLYBC	REAL*8
	L2SOR	LOGICAL*4	•	DUMP	REAL*8
	READ1	LOGICAL*4		INIT2	REAL*8
·	SOR2L	LOGICAL#4		READ1	REAL*8
	VISCOS	LOGICAL*4			
	VSINIT	LOGICAL*4	F2AIF	HST3D	REAL*8
20000		100101111		APLYBC	REAL*8
ERREXE	HST3D	LOGICAL*4		DUMP	REAL*8
	CLOSE	LOGICAL*4	,	INIT2	REAL*8
	ITER	LOGICAL*4		READ1	REAL*8
	SUMCAL	LOGICAL*4			
	TOFEP	LOGICAL*4 LOGICAL*4	F6	ZONPLT	CHARACTER*20
	WBBAL WELLSS	LOGICAL*4 LOGICAL*4		•	
	WELLSS WELRIS	LOGICAL*4	<b>F</b> 7	ZONPLT	CHARACTER*20
		LOGICAL*4			
	WFDYDZ WRITE5	LOGICAL*4	F8	ZONPLT	CHARACTER*20

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
FCJ	WELRIS	REAL*8	FIR	HST3D	REAL*8
				SUHCAL	REAL*8
FDDP	ITER	REAL*8	•	WRITE5	REAL*8
FDSMTH	HST3D	REAL*8	FIRO	HST3D	REAL*8
	APLYBC	REAL*8		APLYBC	REAL*8
	ASEMBL	REAL*8		DUMP	REAL*8
	COEFF	REAL*8		INIT2	REAL*8
	DUMP	REAL*8		READ1	REAL*8
	READ1	REAL*8		SUMCAL	REAL*8
	READ2	REAL*8		WRITE2	REAL*8
	SBCFLO	REAL*8		WRITES	REAL*8
	SUMCAL	REAL*8			
	WBCFLO	REAL*8	FIRN	SUHCAL	REAL*8
	WELLSS	REAL*8			
	WRITE2	REAL*8	FIRVO	hst3D	REAL*8
	WRITES	REAL*8		APLYBC	REAL*8
				DUMP	REAL*8
FDTMTH	HST3D	REAL*8		INIT2	REAL*8
	APLYBC	REAL*8		READ1	REAL*8
	ASEMBL	REAL*8		SUMCAL	REAL*8
	CALCC	REAL*8		WRITE2	REAL*8
	DUMP	REAL*8		WRITE5	REAL*8
	READ1	REAL*8			
	READ2	REAL*8	· FLQ	ZONPLT	REAL*8
	SBCFLO	REAL*8			
	SUHCAL	REAL*8	FL1	ZONPLT	REAL*8
	WBCFLO	REAL*8			
	WELLSS	REAL*8	FL2	ZONPLT	REAL*8
	WRITE2	REAL*8			
	WRITE5	REAL*8	FL3	ZONPĻT	REAL*8
FFPIIL	WFDYDZ	REAL*8	FL4	ZONPLT	REAL*8

Table 11.2--Cross-reference list of variables--Continued

.

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
FL5	ZONPLT	REAL*8	FRESUR	HST3D	LOGICAL*4
			1	ASEMBL	LOGICAL*4
FL6	ZONPLT	REAL*8		CALCC	LOGICAL*4
				CLOSE	LOGICAL*4
FLOREV	WBBAL	LOGICAL*4		COEFF	LOGICAL*4
	WELLSS	LOGICAL*4		DUMP	LOGICAL*4
				ERROR2	LOGICAL*4
FLOW	WELLSS	LOGICAL*4		ETOM1	LOGICAL*4
•		-		INIT2	LOGICAL*4
FMAX	BSODE	REAL*8		INIT3	LOGICAL*4
				IREWI	LOGICAL*4
FMT	ZONPLT	CHARACTER*20		PLOTOC	LOGICAL*4
٦				READ1	LOGICAL*4
FMTL	ZONPLT	CHARACTER*20		READ2	LOGICAL*4
		· .		READ3	LOGICAL*4
FPR3	WELLSS	REAL*8		REWI	LOGICAL*4
			· · ·	SUMCAL	LOGICAL*4
FRAC	COEFF	REAL*8 DIMENSION(*)		WELLSS	LOGICAL*4
	INIT2	REAL*8 DIMENSION(*)		WRITE1	LOGICAL*4
	SUMCAL	REAL*8 DIMENSION(*)		WRITE2	LOGICAL*4
	WELLSS	REAL*8 DIMENSION(*)		WRITE3	LOGICAL*4
	WRITE2	REAL*8 DIMENSION(*)		WRITE5	LOGICAL*4
	WRITE5	REAL*8 DIMENSION(*)			
			FRFAC	WFDYDZ	REAL*8
FRACN	CALCC	REAL*8	FDFIM		004140
•			FRFLM	WELLSS	REAL*8
FRACNP	CALCC	REAL*8			
			FRFLP	WELLSS	REAL*8

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable i type	Variable name	Refer- encing programs	Variable type
FS	INTERP	REAL*8 DIMENSION(*,*)	FTZYDM	CRSDSP	REAL*8
FSCON	INIT2	LOGICAL*4 LOGICAL*4	FTZYDP	CRSDSP	REAL*8
	SUHCAL	LUGICAL-4	GAMMA	APLYBC	REAL*8 DIMENSION(10)
FSLOW	INIT2	LOGICAL*4	<b>UNITA</b>	ALTE.	ALAL-O DIMENSION(IU)
I JLAW	SUMCAL	LOGICAL*4	GCOSTI	HST3D	REAL*8
	DOMONE		0000111	WELRIS	REAL*8
FTDAIF	HST3D	REAL*8		WFDYDZ	REAL*8
	APLYBC	REAL*8			
	DUMP	REAL*8	GRAV	INIT2	REAL*8
	INIT2	REAL*8		•	
	READ1	real*8	GX	HST3D	REAL*8
				APLYBC	REAL*8
FTXYDH	CRSDSP	REAL*8		ASEMBL	REAL*8
				CALCC	REAL*8
FTXYDP	CRSDSP	REAL*8		COEFF	REAL*8
				DUMP	REAL*8
FTXZDH	CRSDSP	REAL*8		ERROR3	REAL*8
				INIT2	REAL*8
FTXZDP	CRSDSP	REAL*8		READ1	REAL*8
				READ3	REAL*8
FTYXDM	CRSDSP	REAL*8		SUHCAL	REAL*8
	enenen	55440		WELLSS	REAL*8
FTYXDP	CRSDSP	REAL*8		WELRIS	REAL*8
FTYZDH	CRSDSP	REAL*8			
FTYZDP	CRSDSP	REAL*8			
FTZXDH	CRSDSP	REAL*8			
FTZXDP	CRSDSP	REAL#8			

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
GY	IIST3D	REAL*8	HEAT	HST3D	LOGICAL*4
	APLYBC	REAL*8		APLYBC	LOGICAL*4
	ASEMBL	REAL*8		ASEMBL	LOGICAL*4
	CALCC	REAL*8		CALCC	LOGICAL*4
	COEFF	REAL*8		COEFF	LOGICAL*4
	DUMP	REAL*8		CRSDSP	LOGICAL*4
	ERROR3	REAL*8		DUMP	LOGICAL*4
	INIT2	REAL*8	-	ERROR2	LOGICAL*4
	READ1	REAL*8		ERROR3	LOGICAL*4
	READ3	REAL*8		ETOM1	LOGICAL*4
	SUMCAL	REAL*8		ETOM2	LOGICAL*4
	WELLSS	REAL*8	,	INIT1	LOGICAL*4
	WELRIS	REAL*8		INIT2	LOGICAL*4
				INIT3	LOGICAL*4
GZ	HST3D	REAL*8		ITER	LOGICAL*4
	APLYBC	REAL*8		PLOTOC	LOGICAL*4
	ASEMBL	REAL*8		READ1	LOGICAL#4
	CALCC	REAL*8		READ2	LOGICAL*4
· •	COEFF	REAL*8		READ3	LOGICAL*4
	DUMP	REAL*8		SUMCAL	LOGICAL*4
	ERROR3	REAL*8		VISCOS	LOGICAL#4
	INIT2	REAL#8		WBBAL	LOGICAL*4
	READI	REAL#8		WELLSS	LOGICAL*4
	READ3	REAL*8		WRITE1	LOGICAL*4
	SUMCAL	REAL#8		WRITE2	LOGICAL*4
	WELLSS	REAL*8		WRITE3	LOGICAL*4
	WELRIS	REAL*8		WRITE4	LOGICAL*4
				WRITE5	LOGICAL*4
HDPRNT	INIT2	REAL*8 DIMENSION(*)			
	SUMCAL	REAL*8 DIMENSION(*)	HTCU	WELRIS	REAL*8
	WRITE2	REAL*8 DIMENSION(*)			
	WRITES	REAL*8 DIMENSION(*)		•	

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
HTCWR	ETOHI	REAL*8 DIMENSION(*)	I	HST3D	INTEGER*4
	READ2	REAL*8 DIMENSION(*)		APLYBC	INTEGER*4
	WELRIS	REAL*8		ASEMBL	INTEGER*4
	WRITE2	REAL*8 DIMENSION(*)		BSODE	INTEGER*4
				COEFF '	INTEGER*4
HWT	ERROR2	REAL*8 DIMENSION(*)		CRSDSP	INTEGER*4
	etom1	REAL*8 DIMENSION(*)		D4DES	INTEGER*4
	INIT2	REAL*8 DIMENSION(*)		DUMP	INTEGER*4
	READ2	REAL*8 DIMENSION(*)		ERROR 1	INTEGER*4
				ERROR2	INTEGER*4
				ERROR3	INTEGER*4
				eton 1	INTEGER*4
				INIT1	INTEGER*4
				INIT2	INTEGER*4
				INIT3	INTEGER*4
	•			INTERP	INTEGER*4
				IREWI	INTEGER*4
				HAP2D	INTEGER*4
				ORDER	INTEGER*4
				PLOT	INTEGER*4
				PLOTOC	INTEGER*4
				PRNTAR	INTEGER*4
				READ1	INTEGER*4
				READ2	INTEGER*4
				READ3	INTEGER*4
				REWI	INTEGER*4
				REWI 3	INTEGER*4
				SUMCAL	INTEGER*4
				TOFEP	INTEGER*4
				VSINIT	INTEGER*4
				WELLSS	INTEGER*4
				WRITE?	INTEGER*4
				WRITE3	INTEGER*4
				WRITE5	INTEGER*4
				ZONPLT	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
I1	DUMP t	INTEGER*4	I21Z	ZONPLT	INTEGER*4 DIMENSION(*)
	IREWI	INTEGER*4		•	
· .	• • • • • • •	INTEGER*4	1222	ZONPLT	INTEGER*4 DIMENSION(*)
	ORDER	INTEGER*4			
	READ1	INTEGER*4	I 2 Z	COEFF	INTEGER*4 DIMENSION(*)
	READ2	INTEGER*4		ERROR2	INTEGER*4 DIMENSION(*)
	READ3	INTEGER*4		INIT2	INTEGER*4 DIMENSION(*)
	REWI	INTEGER*4		READ2	INTEGER*4 DIMENSION(*)
	REWI3	INTEGER*4	*	WRITE2	INTEGER*4 DIMENSION(*)
	SOR2L	INTEGER*4			
	ZONPLT	INTEGER*4	13	ORDER	INTEGER*4
				SOR2L	INTEGER*4
I11Z	ZONPLT	INTEGER*4 DIMENSION(*)		ZONPLT	INTEGER*4
I 12X	PRNTAR	CHARACTER*4	I31Z	ZONPLT	INTEGER*4 DIMENSION(*)
1122	ZONPLT	INTEGER*4 DIMENSION(*)	1322	ZONPLT	INTEGER*4 DIMENSION(*)
112	COEFF	INTEGER*4 DIMENSION(*)	IA1HC	HST3D	INTEGER*4
	ERROR2	INTEGER*4 DIMENSION(*)		DUMP	INTEGER*4
	INIT2	INTEGER*4 DIMENSION(*)		INITI	INTEGER*4
	READ2	INTEGER*4 DIMENSION(*)		READ1	INTEGER*4
	WRITE2	INTEGER*4 DIMENSION(*)			•
			IA2HC	HST3D	INTEGER*4
12	DUMP	INTEGER*4	•	DUMP	INTEGER*4
	IREWI	INTEGER*4		INIT1	INTEGER*4
	MAP2D	INTEGER*4		READ1	INTEGER*4
	ORDER	INTEGER*4		1	, ,
	READ1	INTEGER*4	IA3HC	HST3D	INTEGER*4
	READ2	INTEGER*4	•	DUMP	INTEGER*4
	READ3	INTEGER*4	•	INIT1	INTEGER*4
	REWI	INTEGER*4		READ1	INTEGER*4
	REWI3	INTEGER*4		•	
	SOR2L	INTEGER*4			
	ZONPLT	INTEGER*4	1		

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
IA4	HST3D	INTEGER*4	IABPH	HST3D	INTEGER*4
	DUHP	INTEGER*4		DUMP	INTEGER*4
	INITI	INTEGER*4		INIT1	INTEGER*4
	ITER	INTEGER#4	`	READ1	INTEGER*4
	READ1	INTEGER*4			
			IAIF	HST3D	INTEGER*4
A4B	D4DES	INTEGER*4		APLYBC	INTEGER*4
	-			DUMP	INTEGER*4
AA1	HST3D	INTEGER*4		ERROR2	INTEGER*4
•	DUMP	INTEGER*4		ETOM1	INTEGER*4
	INIT1	INTEGER*4		INIT2	INTEGER*4
	READ1	INTEGER*4		READ1	INTEGER*4
				READ2	INTEGER*4
AA2	HST3D	INTEGER*4		WRITE2	INTEGER*4
	DUMP	INTEGER*4			
	INIT1	INTEGER*4	IALBC	HST3D	INTEGER*4
	READ1	INTEGER*4		DUMP	INTEGER*4
				INIT1	INTEGER*4
AA3	HST3D	INTECER*4		ITER	INTEGER*4
	DUHP	INTEGER*4		READ1	INTEGER*4
	INIT1	INTECER*4			
	READ1	INTEGER*4	IALFL	HST3D	INTEGER*4
				DUMP	INTEGER*4
AA4	HST3D	INTEGER*4		INITI	INTEGER*4
	DUMP	INTEGER*4		INIT2	INTEGER*4
	INIT1	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4			
			IALFT	HST3D	INTEGER*4
AAIF	list3d	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INIT1	INTEGER*4
	INITI	INTEGER*4		INIT2	INTEGER*4
	ITER	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4			

Tabla	11 2 Crass-rafarance	list	~*	variablesContinued
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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
IAMAP	HST3D	INTEGER*4	IAYFBC	HST3D	INTEGER*4
				DUMP	INTEGER*4
IAPRT	IIST3D	INTEGER*4		INIT1	INTEGER*4
			,	READ1	INTEGER*4
IARHC	DUMP	INTEGER*4			
	INIT1	INTEGER*4	IAZFBC	list3D	INTEGER*4
	READ1	INTEGER*4		DUMP	INTEGER*4
			- 0	INIT1	INTEGER*4
IARX .	HST3D	INTEGER*4		READ1	INTEGER*4
	DUMP	INTEGER*4		a. 4	
	INIT1	INTEGER*4	IB	SOR2L	INTEGER*4
	READ1	INTEGER*4	1.1		
·	1	н. -	IBAIF	HST3D	INTEGER*4
IARXBC	HST3D	INTEGER*4		DUMP	INTEGER*4
	н н. • Ма			INIT1	INTEGER*4
IARY	HST3D	INTEGER*4		ITER	INTEGER*4
1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	DUMP	INTEGER*4		READ1	INTEGER*4
	INIT1	INTEGER*4			
	READ1	INTEGER*4	IBBLBC	HST3D	INTEGER*4
				DUMP	INTEGER*4
IARYBÇ	HST3D	INTEGER*4		INITI	INTEGER*4
				ITER	INTEGER*4
IARZ	HST3D	INTEGER*4		READ1	INTEGER*4
	DUMP	INTEGER*4			
· ·	INIT1	INTEGER*4			
	READ1	INTEGER*4			
IARZBC	HST3D	INTEGER*4			
IAXFBC	HST3D	INTEGER*4			
	DUMP	INTEGER*4			
	INIT1	INTEGER*4			
	READ1	INTEGER*4			

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
BC	APLYBC	INTEGER*4 DIMENSION(*)	IC	HST3D	INTEGER*4
	ASEMBL	INTEGER*4 DIMENSION(*)		ASEMBL	INTEGER*4
	COEFF	INTEGER*4 DIMENSION(*)		DUMP	INTEGER*4
	ERROR2	INTEGER*4 DIMENSION(*)		ERROR2	INTEGER*4
	ERROR3	INTEGER*4 DIMENSION(*)		ERROR3	INTEGER*4
	ETOH1	INTEGER*4 DIMENSION(*)		ETOM1	INTEGER*4
	INIT2	INTEGER*4 DIMENSION(*)		INIT1	INTEGER*4
	INIT3	INTEGER*4 DIMENSION(*)		INIT2	INTEGER*4
	ITER	INTEGER*4 DIMENSION(*)		INIT3	INTEGER*4
	ORDER	INTEGER*4 DIMENSION(*)		READ1	INTEGER*4
	READ2	INTEGER*4 DIMENSION(*)		SOR2L	INTEGER*4
	SBCFLO	INTEGER*4 DIMENSION(*)		WRITE3	INTEGER*4
	SOR2L	INTEGER*4 DIMENSION(*)		WRITE5	INTEGER*4
	SUHCAL	INTEGER*4 DIMENSION(*)		ZONPLT	INTEGER*4
	WBBAL	INTEGER*4 DIMENSION(*)			
	WELLSS	INTEGER*4 DIMENSION(*)	ICAIF	HST3D	INTEGER*4
	WRITE2	INTEGER*4 DIMENSION(*)		DUMP	INTEGER*4
	WRITE3	INTEGER*4 DIMENSION(*)			
	WRITE4	INTEGER*4 DIMENSION(*)		INITI	INTEGER*4
	WRITE5	INTEGER*4 DIMENSION(*)		ITER	INTEGER*4
				READ1	INTEGER*4
IBCMAP	WRITE5	INTEGER*4 DIMENSION(*)			
			ICALL	IREWI	INTEGER*4
IBLBC	HST3D	Integer*4		REWI	INTEGER*4
	DUMP	INTEGER*4		REWI3	INTEGER*4
	INIT1	INTEGER*4			
	ITER	INTEGER*4	ICC	HST3D	LOGICAL*4
	READ1	INTEGER*4		DUMP	LOGICAL*4
				ERROR2	LOGICAL*4
				ETOM1	LOGICAL*4
				INIT2	LOGICAL*4
				READI	LOGICAL*4
				READ2	LOGICAL*4
			(	WRITE2	LOGICAL*4

Table 11 2- Charte materia	12.4		
Table 11.2Cross-reference	list of	variablesContinued	

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Variable name	Refer- encing programs	Variable type	Variable nam <del>e</del>	Refer- encing programs	Variable type
ICC24	HST3D	INTEGER*4	ICHYDP	HST3D	LOGICAL*4
	DUMP	INTEGER#4		DUMP	LOGICAL*4
	INITI	INTEGER*4		ERROR2	LOGICAL*4
	ITER	INTEGER*4		ETOM1	LOGICAL*4
	READ1	INTEGER*4		INIT2	LOGICAL*4
				READ1	LOGICAL*4
ICC34	HST3D	INTEGER*4	5	READ2	LOGICAL*4
	DUMP	INTEGER*4	• .	WRITE2	LOGICAL*4
• ,	INIT1	INTEGER*4			
	ITER	INTEGER*4	ICI	HST3D	INTEGER*4
	READ1	INTEGER*4		DUMP	INTEGER*4
	-	· · · · · · · · · · · · · · · · · · ·		INITI	INTEGER*4
ICC35	HST3D	INTEGER*4		ITER	INTEGER*4
,	DUMP	INTEGER*4		READ1	INTEGER*4
	INIT1	INTEGER*4		• •	
1	ITER	INTEGER*4	ICLBC	HST3D	INTEGER*4
1.1	READ1	INTEGER*4		DUMP	INTEGER*4
				INIT1	INTEGER*4
ICCW	HST3D	INTEGER*4	,	ITER	INTEGER*4
1	12-AV		r 1	READ1	INTEGER*4
ICFLX	HST3D	INTEGER*4			·
	DUMP	INTEGER*4	ICMAX	HST3D	INTEGER*4
	INIT1	INTEGER*4		SUMCAL	INTEGER*4
	ITER	INTEGER*4		WRITES	INTEGER*4
	READ1	INTEGER*4			
-	e		ICNP	HST3D	INTEGER*4
ICH	ZONPLT	INTEGER*4			
ICHWT	HST3D	LOGICAL*4	ICON	D4DES	INTEGER*4
	DUMP	LOGICAL*4			• .
	ERROR2	LOGICAL*4	ICOW	HST3D	INTEGER*4
	ETOM1	LOGICAL*4			
	INIT2	LOGICAL*4	ICQFLX	HST3D	INTEGER*4
	READ1	LOGICAL*4			
	READ2	LOGICAL*4			
	WRITE2	LOGICAL*4			

Table 11.2--Cross-reference list of variables--Continued

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	Refer-			Refer-	
Variable	encing	Variable	Variable	encing	Variable
name	programs	type	name	programs	type
ICSBC	HST3D	INTEGER*4	ID	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INIT1	Integer*4		INITI	INTEGER*4
	ITER	INTEGER*4		L2SOR	Integer*4
	READ1	INTEGER*4		READ1	Integer*4
ICT	HST3D	LOGICAL*4	ID4	ORDER	INTEGER*4
	DUMP	LOGICAL*4			
	ERROR2	LOGICAL*4	ID4CON	D4DES	INTEGER*4 DIMENSION(6)
	ETOH1	LOGICAL*4			
	INIT2	LOGICAL*4	I D4NO	ASEMBL	INTEGER*4 DIMENSION(*)
	READ1	LOGICAL#4		D4DES	INTEGER*4 DIMENSION (*)
	READ2	LOGICAL*4		ORDER	INTEGER*4 DIMENSION (*)
	WRITE2	LOGICAL*4		SBCFLO	INTEGER*4 DIMENSION (*)
				WBCFLO	INTEGER*4 DIMENSION (*)
I CWKT	HST3D	Integer*4			
	DUMP	Integer*4	IDBKD	list3d	INTEGER*4
	INIT1	INTEGER*4		DUHP	INTEGER*4
	ITER	Integer*4		INITI	Integer*4
	READ1	INTEGER*4		ITER	INTEGER*4
	SUMCAL	INTEGER*4		READ1	Integer*4
ICXH	ASEMBL	INTEGER*4	IDC	HST3D	INTEGER*4
				DUMP	INTEGER*4
ICXP	ASEMBL	INTECER*4		INIT1	INTEGER*4
				READ1	INTEGER*4
ICYM	ASEHBL	Integer*4			
			IDEN	HST3D	INTEGER*4
ICYP	ASEMBL	INTEGER*4		DUMP	INTECER*4
				INIT1	INTEGER*4
ICZM	ASEMBL	INTEGER*4		ITER	INTEGER*4
				READ1	INTEGER*4
ICZP	ASEHBL	INTEGER*4			
			IDENW	HST3D	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
IDIAG	D4DES	INTEGER*4 DIMENSION(*)	IDPWKT	HST3D	INTEGER*4
	ORDER	INTEGER*4 DIMENSION(*)		DUMP	INTEGER*4
				INIT1	INTEGER*4
DIR	L2SOR	INTEGER*4		ITER	INTEGER*4
	ORDER	INTEGER*4	.: •	READ1	INTEGER*4
	SOR2L	INTEGER*4		SUMCAL	INTEGER*4
DNFBC	HST3D	INTEGER*4	IDQHDT	HST3D	INTEGER*4
	DUMP	INTEGER*4	-	DUMP	INTEGER*4
	INITI	INTEGER*4		INITI	INTEGER*4
	ITER	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4		· · · ·	
			IDQWDP	HST3D	INTEGER*4
DNLBC	HST3D	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INITI	INTEGER*4
	INIT1	INTEGER*4		ITER	INTEGER*4
	ITER	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4		SUMCAL	INTEGER*4
DNOAR	HST3D	INTEGER*4	IDRCON	D4DES	INTEGER*4 DIMENSION(6)
	DUMP	INTEGER*4			
	INIT1	INTEGER*4	IDŤ	HST3D	INTEGER*4
	ITER	INTEGER*4		DUMP	INTEGER*4
1	READ1	INTEGER*4		INITI	INTEGER*4
				READ1	INTEGER*4
IDNUM	PLOT	INTEGER*4			· ·
			IDTHIC	HST3D	INTEGER*4
IDP	HST3D	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INIT1	INTEGER*4
	INIT1	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4			

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variabl <i>e</i> type	Variable name	Refer- encing programs	Variable type
IDTHWR HST3D	HST3D	INTEGER*4	IENDIV	list3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	IN <b>IT1</b>	INTEGER*4		ERROR 1	INTECER*4
	INIT2	INTEGER*4		INITI	INTEGER*4
	READ1	Integer*4		READ1	INTEGER*4
	WBBAL	INTEGER*4		WRITE1	INTEGER*4
	WELLSS	Integer*4			
			IENDVV	HST3D	INTEGER*4
IE	ERRPRT	INTEGER*4		DUMP	INTEGER*4
				ERROR 1	INTEGER*4
IE1	ERRPRT	INTEGER*4		INITI	INTEGER*4
				READ1	INTEGER*4
E2	ERRPRT	INTEGER*4		WRITEI	INTEGER*4
IEN	list3D	INTEGER*4	IEQ	HST3D	INTEGER*4
	DUMP	INTEGER*4	•	ASEHBL	INTEGER*4
	INIT1	Integer*4		CALCC	INTEGER*4
	ITER	INTEGER*4		ITER	INTEGER*4
	READ1	Integer*4		L2SOR	INTEGER*4
				SBCFLO	INTEGER*4
IEHSUR	HST3D	INTEGER*4		SOR2L	INTEGER*4
	DUMP	INTEGER*4			
	INIT1	INTEGER*4	IER	IREWI	INTEGER*4
	ITER	INTEGER*4		REWI	INTEGER*4
	READ1	INTEGER*4		REWI3	INTEGER*4
	SUHCAL	INTEGER*4			
IEHWKT	list3D	INTEGER*4			
	DUMP	INTEGER*4			
	INITI	INTEGER*4			
	ITER	INTEGER*4			
	READ1	INTEGER*4			
		-venon-L/			

Table 11.2--Cross-reference list of variables--Continued

SUHCAL

INTEGER\*4

Jariable	Refer- encing	Variable	Variable	Refer- encing	Variable
name	programs	type	name	programs	type
ERR	HST3D	LOGICAL*4 DIMENSION(200)	INTCR	HST3D	INTEGER#4
	ERROR 1	LOGICAL*4 DIMENSION(200)		DUMP	INTEGER*4
	ERROR2	LOGICAL*4 DIMENSION(200)		INIT1	INTEGER*4
	ERROR3	LOGICAL*4 DIMENSION(200)		INIT2	INTEGER*4
	ERRPRT	LOGICAL*4 DIMENSION(200)		READ1	INTEGER*4
	INIT2	LOGICAL*4 DIMENSION(200)		WBBAL	INTEGER*4
	IREWI	LOGICAL*4 DIMENSION(200)		WELLSS	INTEGER*4
	ITER	LOGICAL*4 DIMENSION(200)			
	READ1	LOGICAL*4 DIMENSION(200)	IHWT	HST3D	INTEGER*4
	READ2	LOGICAL*4 DIMENSION(200)	_	•	
	READ3	LOGICAL*4 DIMENSION(200)	II	ASEMBL	INTEGER*4
	REWI	LOGICAL*4 DIMENSION(200)		MAP2D	INTEGER*4
	REWI3	LOGICAL*4 DIMENSION(200)			
	SOR2L	LOGICAL*4 DIMENSION(200)	III	SOR2L	INTEGER*4
	SUMCAL	LOGICAL*4 DIMENSION(200)		-	
			II12	PRNTAR	CHARACTER*4
IF12	PRNTAŘ	CHARACTER*4			
			II1P1	SOR2L	INTEGER*4
IFACE	IREWI	INTEGER*4			1
		•	IIIZ	HST3D	INTEGER*4
IFMT	PRNTAR	INTEGER*4		DUMP	INTEGER*4
	WRITE2	INTEGER*4		INIT1	INTEGER*4
				READ1	INTEGER*4
IFORM	PRNTAR	CHARACTER*84			
			1122	HST3D	INTEGER*4
IFRAC	HST3D	INTEGER*4		DUMP	INTEGER*4
				INITI	INTEGER*4
IG12	PRNTAR	CHARACTER*6		READ1	INTEGER*4
IHDPRT	HST3D	INTEGER*4	113	SOR2L	INTEGER*4
			II32	ZONPLT	INTEGER*4
		•	115	PRNTAR	CHARACTER*6

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	Refer-			Refer-	
Variable	encing	Variable	Variable	encing	Variable
name	programs	type	name	programs	type
11BC	HST3D	INTEGER*4	IJ2Z	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INIT1	INTEGER*4		INIT1	INTEGER*4
	L2SOR	INTEGER*4		READ 1	INTEGER*4
	READ1	INTEGER*4			
			IJW	list3D	INTEGER*4
IIBCMP	HST3D	Integer*4		DUMP	INTEGER*4
				INIT1	INTEGER*4
1104	HST3D	INTEGER*4		ITER	INTEGER*4
	DUMP	INTEGER*4		READ1	INTEGER*4
	INIT1	INTEGER*4		SUHCAL	INTEGER*4
	ITER	Integer*4		•	
	READ1	INTEGER*4	IJWEL	HST3D	INTEGER*4
IIDAG	HST3D	INTEGER*4	IK1Z	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INIT1	INTEGER*4		INIT1	INTEGER*4
	ITER	INTEGER*4		READ1	INTEGER*4
	READ1	INTECER*4			
			IK2Z	HST3D	INTEGER*4
IIFMT	PRNTAR	Intecer*4		DUMP	INTEGER*4
	•			INIT1	INTEGER*4
IINZON	HST3D	INTEGER*4		READ1	INTEGER*4
			IKARHC	HST3D	INTEGER*4
IIR2	PRNTAR	INTEGER*4			
			IKLBC	HST3D	INTECER*4
IIW	HST3D	INTEGER*4		DUHP	INTEGER*4
	DUMP	INTEGER*4		INIT1	INTEGER*4
	INIT1	INTEGER*4		ITER	INTEGER*4
	ITER	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4			
	SUHCAL	INTEGER*4			
IJIZ	HST3D	INTEGER*4			
	DUMP	INTEGER*4	1		
	INITI	INTEGER*4			
	READ1	INTEGER*4	(		

Variable	Refer- encing	Variable	Variable	Refer- encing	Variable
name	programs	type	name	programs	type
IKTAWR	HST3D	INTEGER*4	IKTZPH	HST3D	INTEGER*4
•	DUMP	INTEGER*4			· .
	INIT1	INTEGER*4	IKXX	HST3D	INTEGER*4
	INIT2	INTEGER*4		DUMP	INTEGER*4
	READ1	INTEGER*4		INITI	INTEGER*4
	WBBAL	INTEGER*4		READ1	INTEGER*4
1	WELLSS	INTEGER*4			
			IKYY	HST3D	INTEGER*4
IKTHX	HST3D	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INIT1	INTEGER*4
	INIT1	INTEGER*4		READI	INTEGER*4
	READ1	INTEGER*4			
	·		IKZZ	hst3d	INTEGER*4
IKTHY	HST3D	INTEGER*4	· · ·	DUMP	INTEGER*4
	DUMP	INTEGER*4		INITI	INTEGER*4
	INITI	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4			
		• •	ILAVFL	HST3D	INTEGER*4
IKTHZ	HST3D	INTEGER*4		DUMP	INTEGER*4
191	DUMP	INTEGER*4		ERROR1	INTEGER*4
	INIT1	INTEGER*4		INITI	INTEGER*4
	READ1	INTEGER*4		READI	INTEGER*4
		•		WRITE1	INTEGER*4
IKTWR	HST3D	INTEGER*4			
•	DUMP	INTEGER*4	ILBL	PLOT	INTEGER*4
	INIT1	INTEGER*4			
	INIT2	INTEGER*4	ILBW	INITI	INTEGER*4
•.	READ1	INTEGER*4			
	WBBAL	INTEGER*4	ILCBW	HST3D	INTEGER*4
	WELLSS	INTEGER*4		DUMP	INTEGER*4
				INIT1	INTEGER*4
IKTXPM	HST3D	INTEGER*4		ITER	INTEGER*4
				READI	INTEGER*4
ІКТҮРМ	HST3D	INTEGER*4	1	SUMCAL	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
ILCTW	HST3D	INTEGER#4	Інар	READ3	INTEGER*4
	DUMP	INTEGER#4			· · ·
	INIT1	INTEGER*4	IMAP1	HST3D	INTEGER*4 DIMENSION(3)
	ITER	INTEGER*4		ERROR3	INTEGER*4 DIMENSION(3)
	READ1	Integer#4		ETOM2	INTEGER+4 DIMENSION(3)
	SUMCAL	Integer#4		INIT3	INTEGER*4 DIMENSION(3)
				READ3	INTEGER*4 DIMENSION(3)
ILH	D4DES	Intecer*4		WRITE3	INTEGER*4 DIMENSION(3)
				WRITE5	INTEGER*4 DIMENSION(3)
ILIVPA	HST3D	INTEGER*4			
	DUMP	INTEGER*4	IHAP2	HST3D	INTEGER*4 DIMENSION(3)
	ERROR1	INTEGER#4		ERROR3	INTEGER*4 DIMENSION(3)
	INITI	INTEGER*4		ETOH2	INTEGER*4 DIMENSION(3)
	READ1	INTEGER*4		INIT3	INTEGER*4 DIMENSION(3)
	WRITE!	INTEGER=4		READ3	INTEGER*4 DIMENSION(3)
-				WRITE3	INTEGER*4 DIMENSION(3)
ILNXI	WELRIS	REAL*8		WRITES	INTEGER*4 DIMENSION(3)
ILPRT	HST3D	INTEGER*4	IMAX	ORDER	INTEGER*4
	DUMP	INTEGER#4			
	init1	INTECER#4	IMFBC	HST3D	INTEGER*4
	READI	INTEGER#4		DUMP	INTEGER*4
				INITI	INTEGER#4
ILVPA	HST3D	INTEGER#4		ITER	INTEGER*4
	DUMP	INTEGER*4		READ1	INTEGER*4
	ERROR1	INTEGER*4			
	INIT1	INTEGER*4	IMICBC	HST3D	INTEGER*4
	READ1	INTEGER*4		DUMP	INTEGER*4
	WRITEL	INTEGER*4		INIT1	INTEGER*4
				ITER	INTEGER*4
IMAIFC	HST3D	INTEGER+4		READ1	INTEGER*4
	DUMP	INTEGER*4			
	INIT1	INTEGER*4	IMIN	ORDER	INTEGER*4
	ITER	INTEGER*4	266217		
	READ1	INTEGER*4			

Table 11.2--Cross-reference list of variables--Continued

	Refer-			Refer-	
Variable	encing	Variable	Variable	encing	Variable
name	programs	type	name	programs	type
IMLBC	HST3D	INTEGER*4	INCI	READ2	INTEGER*4
	DUMP	Integer*4		READ3	Integer*4
	INITI	INTEGER*4			
	ITER -	INTEGER*4	INCJ	READ2	INTEGER*4
	READ1	INTEGER*4		READ3	Integer*4
Imm	INIT2	INTEGER*4	INCJI	SOR2L	INTEGER*4
IMOBW	HST3D	INTEGER*4	INCJ2	SOR2L	INTEGER*4
Imod	REWI	INTEGER*4	INCJ3	SOR2L	INTEGER*4
	REWI3	INTEGER*4 DIMENSION(3)			
			Inzone	ERROR2	DIMENSION(*) LOGICAL*
Impptc	WRITE3	INTEGER*4			
	WRITE5	INTEGER*4	IOR	PRNTAR	INTEGER*4
IMPQW	HST3D	LOGICAL*4	IOUT	HAP2D	INTEGER*4
1 <sup>-</sup>	DUMP	LOGICAL#4			
	etomi	logical#4	IP	HST3D	INTEGER*4
	INIT2	logical#4		DUMP	INTEGER#4
	ITER	LOGICAL*4		INIT1	Integer+4
	READ1	LOGICAL*4		READ1	Integer*4
	READ2	LOGICAL*4		WRITE5	INTEGER*4
	WBBAL	LOGICAL*4			
	WELLSS	LOGICAL*4	IPAGE	MAP2D	Integer*4
•	WELRIS	LOGICAL#4			
	WRITE2	LOGICAL*4	IPAIF	HST3D	INTEGER*4
	WRITE5	LOGICAL*4		DUMP	INTEGER*4
				INIT1	INTEGER*4
IMSBC	HST3D	INTEGER*4		ITER	INTEGER*4
•	DUMP	INTEGER*4		READI	INTEGER*4
	INIT1	INTEGER#4	· · · ·		
	ITER	INTEGER*4			
	READ1	INTEGER*4			

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Table 11.2--Cross-reference list of variables--Continued

ariable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
PAR	IREWI	INTEGER*4 DIMENSION(*)	IPMHV	HST3D	INTEGER#4
	,	·		ASEMBL	Integer#4
PARI	IREWI	INTEGER*4		DUMP	INTEGER#4
				Initi	Integer+4
PCS	HST3D	INTEGER#4	,	READ1	Integer*4
PCW	HST3D	INTEGER#4	IPMZ	COEFF	INTEGER=4
				ERROR2	INTEGER#4
PHILB	HST3D	Integer+4		<b>ETOH1</b>	INTEGER#4
	DUMP	Intecer*4		init2	INTEGER#4
	Init1	INTEGER*4		READ2	INTEGER#4
	READ1	Integer#4		ZONPLT	INTEGER*4
PLOT	MAP2D	INTEGER*4 DIMENSION(*,*)	IPHZ1	ZONPLT	Intecer*4
PHAX	HST3D	INTECER#4	IPMZ2	ZONPLT	INTEGER*4
	SUMCAL	INTEGER#4			
	WRITE5	INTEGER*4	IPNP	HST3D	INTECER#4
рнсну	HST3D	INTEGER#4	IPOR	HST3D	INTECER*4
	ASEMBL	INTEGER*4		DUMP	Integer*4
	DUNP	INTEGER#4		Initl	INTEGER#4
	INITI	INTEGER#4		READ1	INTEGER#4
	READI	INTEGER#4			
			IPOS	HST3D	INTEGER#4
PHCV	HST3D	INTEGER*4			
	ASEMBL	INTEGER+4	IPOW	hst3d	INTEGER#4
	DUMP	INTEGER*4			
	INITI	INTEGER*4			
	READ1	INTEGER*4			
PHCVK	HST3D	INTEGER*4			
	ASEMBL	INTEGER*4			
	DUMP	INTEGER*4			
	INITI	INTEGER*4			
	READI	INTEGER*4			

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable Lype
					· · · · · · · · · · · · · · · · · · ·
IPRPTC	HST3D	INTEGER*4	IPWKT	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INIT2	INTEGER*4		INIT1	INTEGER*4
	READI	INTEGER*4		ITER	INTEGER*4
•	READ2	INTEGER*4		READ ]	INTEGER*4
	READ3	INTEGER*4		SUMCAL	Integer*4
	SUMCAL	INTEGER*4	IPWKTS	HST3D	INTEGER*4
	WRITE2	INTEGER*4	IPWKID		
	WRITE3	INTEGER#4		DUMP INITI	INTEGER*4 INTEGER*4
	WRITE4	INTEGER*4		ITER	INTEGER*4
	WRITE5	INTEGER*4		READ1	
TRADA				SUMCAL	INTEGER*4 INTEGER*4
IPSBC	HST3D	INTEGER*4		DOUCAL	TUTERCK-4
	DUMP	INTEGER*4	IPWSRS	HST3D	INTEGER*4
	INIT]	INTEGER#4	TEMORO	DUMP	INTEGER*4
	ITER READI	INTEGER#4		INITI	INTEGER*4
	KENUL	INTEGER*4	1 - A	ITER	INTEGER*4
IPT	PLOTOC	INTEGER*4		READI	INTEGER*4
AE 4	FROTOC	INLEGER-4		SUMCAL	INTEGER*4
IPV	HST3D	INTEGER*4		DUNGAN	
AL V	ASEMBL	INTEGER*4	IPWSUR	HST3D	INTEGER*4
	DUMP	INTEGER*4	AL HOUL	DUMP	INTEGER*4
	INITI	INTEGER*4		INITI	INTEGER*4
	READI	INTEGER*4		ITER	INTEGER*4
	KEAD1	ANALGER"4		READI	INTEGER*4
IPVK	HST3D	INTEGER*4		SUMCAL	INTEGER*4
TT AU	ASEMBL	INTEGER*4		U VI I VI I VI I I I	6116443MIX ***
	DUMP	INTEGER*4	IQFAIF	HST3D	INTEGER*4
	INITI	INTEGER*4	TALUTE	DUMP	INTEGER*4
	READI	INTEGER*4		INITI	INTEGER*4
,	********			ITER	INTEGER*4
				READ1	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable . type	Variable name	Refer- encing programs	Variable type
IQFBCV	HST3D	INTEGER*4	IQHFX	HST3D	INTEGER*4
	DUMP	INTEGER*4		_	
	in <b>iti</b>	INTEGER=4	IQHPY	HST3D	INTEGER*4
	READ1	Integer#4			
			IQHFZ	HST3D	INTEGER*4
Iqffx	HST3D	INTEGER*4			
			IQHLBC	HST3D	INTEGER*4
Iqffy	HST3D	INTECER*4		DUMP	INTECER*4
				INITL	INTEGER*4
IQFFZ	HST3D	Intecer*4		READ1	INTEGER*4
IQFLBC	HST3D	INTEGER*4	IQHLYR	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INITI	INTECER*4		init1	INTEGER*4
	READ1	INTEGER*4		ITER	INTEGER#4
				READ1	INTEGER*4
1qpsbc	hst3d	Intecer*4		SUMCAL	Integer*4
	DUMP	Integer*4			
	initi	INTEGER*4	Iqhsbc	HST3D	INTEGER#4
	1TER	INTEGER*4		DUMP	INTEGER#4
	READ1	INTEGER*4		INITI	INTEGER*4
				ITER	INTEGER*4
IQHAIF	HST3D	INTEGER*4		READ1	INTEGER*4
	DUMP	INTEGER*4			
	INITI	INTEGER*4	IQHW	HST3D	INTEGER#4
	READ1	INTECER*4		DUMP	INTEGER*4
TANADA	1100055			INIT1	INTECER*4
Iquebe	HST3D	INTEGER#4		ITER	INTEGER#4
	DUMP	INTEGER*4		READ1	INTEGER*4
	INITI	INTEGER*4		SUHCAL	INTEGER*4
	READI	Integer*4			
	110000-		IQSAIF	HST3D	INTEGER*4
lqhfbc	HST3D	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INITI	INTEGER*4
	INITI	INTEGER*4		READ1	INTEGER*4
	ITER	INTEGER*4			
	READ1	INTEGER#4			

	Refer-	•		Refer-	
Variable	encing	Variable	Variable	encing	Variable
пале 🦂	programs	type	name	programs	type
IQSFBC	HST3D	INTEGER*4	IQWLYR	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	initi	INTEGER*4		INITI	INTEGER*4
	ITER	INTEGER*4		ITER	INTEGER*4
	READ]	INTEGER*4		READI	INTEGER*4
				SUMCAL	INTEGER*4
IQSFX	HST3D	INTEGER*4	·		
	· .		IQWM	HST3D	INTEGER*4
IQSFY	HST3D	INTEGER#4		DUMP	INTEGER*4
				INITI	INTEGER*4
IQSFZ	HST3D	INTEGER*4	1	iter	INTEGER*4
				READI	INTEGER*4
IQSLBC	HST3D	Integer*4		SUMCAL	INTEGER*4
•	DUMP	INTEGER*4			
	INITI	Integer*4	IQWV	HST3D	INTEGER*4
	READ1	INTEGER#4		DUMP	INTEGER*4
				INITI	INTEGER*4
IQSLYR	HST3D	INTEGER*4		ITER	Integer*4
•	DUMP	INTEGER*4		READ1	Integer*4
	INITI	INTEGER*4		SUMCAL	Integer*4
	ITER	INTEGER*4			
	READ1	INTEGER*4	IR2	PRNTAR	Integer*4
	SUMCAL	INTEGER*4			
		• .	IR3	PRNTAR	INTEGER*4
IQSSBC	HST3D	INTEGER*4		•	•
	DUMP	INTEGER*4	IRJLBL	PRNTAR	CHARACTER*1
	INITI	INTEGER*4	`		
	ITER	INTEGER*4	IR3P	PRNTAR	INTEGER*4
· ·	READ1	Integer*4			
	t i		IRBW	HST3D	INTEGER*4
IQSW	HST3D	INTEGER*4		DUMP	INTEGER*4
•	DUMP	INTEGER*4		INIT1	INTEGER*4
	INITI	INTEGER*4		itėr	INTEGER*4
	ITER	INTEGER#4		READ1	INTEGER*4
	READ1	INTEGER*4			
	SUMCAL	INTEGER*4			

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable
	hroligne	cype		programs	type
IRCPPM	HST3D	* INTECER#4	IRHSW	IIST3D	INTECER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INI <b>TI</b>	INTEGER*4		INITI	INTEGER*4
	READ1	Integer*4		ITER	INTEGER*4
				READ1	INTEGER*4
IRF	HST3D	Integer#4		SUMCAL	INTEGER*4
	DUMP	Integer#4			
	INITI	INTECER#4	IRM	HST3D	INTEGER*4
	READ1	Integer*4		DUMP	1nteger*4
				init1	INTEGER*4
IRH	HST3D	INTECER*4		READ1	INTEGER*4
	DUMP	INTEGER*4			
	INITI	INTEGER*4	IROW	D4DES	INTEGER*4
	READ1	INTEGER*4			
			IRS	HST3D	INTEGER*4
IRHI	HST3D	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INIT1	INTEGER*4
	INITI	INTEGER*4		READ1	INTEGER*4
	ITER	INTECER#4		•	
	READ1	INTEGER*4	IRSI	HST3D	INTEGER*4
				DUMP	INTEGER*4
IRHS	HST3D	INTEGER*4		INITI	INTEGER#4
	DUMP	INTEGER*4		ITER	INTEGER#4
	INITI	INTEGER*4		READ1	Integer*4
	READ1	INTEGER*4			
			ISIGN	HST3D	INTEGER*4
IRHSBC	HST3D	INTEGER*4		WBBAL	INTEGER*4
	DUMP	INTEGER*4		WELLSS	INTEGER*4
	INITI	INTEGER*4		WELRIS	INTEGER*4
	ITER	INTEGER*4			
	READ1	INTEGER*4	ISORD	HST3D	INTEGER*4 DIMENSION(3)
				DUHP	INTEGER*4 DIMENSION(3
				L2SOR	INTEGER*4 DIMENSION(3)
				READ1	INTEGER*4 DIMENSION(3
				WRITE5	INTEGER*4 DIMENSION(3

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
ISUM	ORDER	INTEGER*4	ITBWR	HST3D	INTEGER*4
				DUMP	INTEGER*4
1SXX	HST3D	INTEGER#4		INITI	INTEGER*4
	DUMP	INTEGER*4		INIT2	INTEGER*4
	INITI	INTEGER*4		READ1	INTEGER*4
	ITER	INTEGER*4		WBBAL	INTEGER*4
	READ1	INTEGER*4		WELLSS	INTEGER*4
ISYY	HST3D	INTEGER*4	ITC	HST3D	INTEGER*4
	DUMP	INTEGER*4			· ·
	INITI	INTEGER*4	ITCS	HST3D	INTEGER*4
	ITER	INTEGER*4			
	READI	INTEGER*4	ITCW	HST3D	INTEGER*4
ISZZ	HST3D	INTEGER*4	ITFLX	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INIT1	INTEGER*4		INIT1	INTEGER*4
	ITER	INTEGER#4		ITER	INTEGER*4
	READ1	INTEGER*4		READ1	INTEGER*4
IT	HST3D	INTEGER#4	ITFW	HST3D	INTEGER*4
	DUMP	INTEGER#4		DUMP	INTEGER#4
	initi	INTEGER*4		INITI	INTEGER*4
	READ1	INTEGER#4		ITER	INTEGER*4
				READ1	INTEGER*4
ITAIF	HST3D	INTEGER*4		SUMCAL	INTEGER*4
	DUMP	INTEGER*4			
	INITI	INTEGER*4	ITPX	HST3D	INTEGER*4
	ITER	INTEGER*4	~~~~~~~	ASEMBL	INTEGER*4
	READ 1	Integer*4		DUMP	INTEGER*4
			•	INITI	INTEGER*4
	·			READ1	INTEGER*4

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
ITFY	HST3D	INTECER+4	ITHY	HST3D	INTEGER*4
	ASEMBL	INTECER+4		ASEMBL	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INITI	INTEGER*4		INIT1	INTEGER*4
	READI	INTEGER*4		READ1	Integer*4
ITFZ	HST3D	INTEGER*4	ITHYX	HST3D	INTEGER*4
	ASEMBL	Intecer*4		ASEMBL	Integer*4
	DUMP	INTEGER#4		DUMP	INTEGER*4
	INIT1	INTEGER*4		INITI	INTECER#4
	READI	INTEGER*4		READI	INTEGER*4
LTHCBC	HST3D	INTEGER*4	ITHYZ	HST3D	INTEGER*4
	DUMP	INTEGER*4		ASEMBL	INTECER*4
	init1	INTEGER*4		DUMP	INTEGER*4
	READI	Intecer*4		INITI	INTEGER*4
				READI	INTEGER*4
LTHX	hst3d	Integer*4			
	ASEMBL	INTEGER*4	ITHZ	HST3D	INTEGER*4
	DUMP	Integer#4		ASEMBL	INTEGER*4
	init1	INTEGER*4		DUMP	INTEGER*4
	READ1	INTEGER*4		INITI	INTEGER*4
	•			READI	Integer+4
ITHXY	hst3d	INTEGER*4			
	ASĖMBL	INTEGER*4	ITHZX	hst3d	INTEGER*4
	DUMP	INTEGER*4		ASEMBL	INTEGER*4
	INITI	INTEGER*4		DUMP	INTECER*4
	READI	INTEGER*4		INITI	INTEGER*4
				READ1	INTEGER*4
ITHXZ	HST3D	INTEGER*4			
	ASEMBL	INTEGER*4	I THZY	HST3D	INTEGER*4
	DUMP	INTEGER*4		ASEMBL	INTEGER*4
	INITI	INTEGER*4		DUMP	INTEGER*4
	READ1	INTEGER*4		'INITI	INTEGER*4
			C	READ1	INTEGER*4

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
LTIME HST31	HST3D	INTEGER*4	ITM2	HST3D	INTEGER*4
	APLYBC	INTEGER*4		DUMP	INTEGER*4
	CALCC	INTEGER*4		INITI	INTEGER*4
	CLOSE	Integer*4	•	L2SOR	INTEGER*4
	COEFF	INTEGER*4		READ1	INTEGER*4
	DUMP	INTEGER*4			
	ERROR3	INTEGER*4	ITMAX	HST3D	Integer*4
	ETOM2	INTEGER*4		SUMCAL	INTEGER*4
	INIT2	INTEGER*4		TOFEP	Integer*4
	INIT3	INTEGER*4		WRITE5	INTEGER*4
	ITER	INTEGER*4		•	
•	READ1	INTEGER*4	ITNO	SOR2L	INTEGER*4
	READ3	INTEGER*4	1		
•	SUMCAL	INTEGER*4	ITNOC	HST3D	INTEGER*4
	WBBAL	INTEGER*4		SOR2L	INTEGER*4
	WELLSS	INTEGER*4	•	WRITE5	INTEGER*4
	WRITE2	INTEGER*4			
	WRITE3	INTEGER*4	ITNOP	HST3D	Integer*4
	WRITE4	INTEGER*4		SOR2L	INTEGER#4
	WRITE5	INTEGER*4		WRITE5	INTEGER*4
ITLBC	HST3D	INTEGER*4	ITNOT	HST3D	INTEGER*4
	DUMP	INTEGER*4		SOR2L	INTEGER*4
	INITI	INTEGER*4		WRITES	INTEGER*4
	ITER	INTEGER*4			
	READ1	INTEGER*4	ITNP	HST3D	INTEGER*4
ITMI	HST3D	INTEGER*4	ITO	HST3D	INTEGER*4
	DUMP	INTEGER*4			
	INITI	integer*4	ITOS	HST3D	Integer*4
	L2SOR	INTEGER*4			
	READI	INTEGER*4	ITOW	HST3D	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
ITPI	HST3D	INTEGER#4	ITSBC	HST3D	INTEGER*4
	DUMP	INTEGER+4		DUMP	INTEGER*4
	init1	Integer#4		INITI	INTEGER*4
	L2SOR	INTECER#4		ITER	INTEGER#4
	READ1	INTEGER#4		READI	INTEGER#4
ITP2	HST3D	INTEGER#4	ITSX	HST3D	INTEGER#4
	DUMP	INTEGER#4		ASEHBL	Integer#4
	initi	INTECER#4		DUMP	INTEGER#4
	L2SOR	INTEGER#4		INIT1	INTEGER*4
	READ1	INTECER#4		READ1	Intecer+4
Ітрнвс	HST3D	INTECER#4	ITSXY	HST3D	INTEGER#4
	DUMP	INTECER#4		ASEMBL	INTEGER#4
	INITI	INTEGER#4		DUMP	INTEGER#4
	READI	INTEGER*4		init1	Intecer+4
				READ1	INTEGER#4
LTQFLX	HST3D	Integer#4			
-			ITSIZ	HST3D	INTEGER+4
ITRN	HST3D	Intecer#4		ASEMBL	INTEGER#4
	ASEMBL	Integer#4		DUMP	INTEGER#4
	ITER	Integer*4		initi	INTEGER*4
	WRITE5	Integer+4		READ1	INTEGER#4
ITRNI	WELLSS	INTEGER#4	ITSY	HST3D	INTEGER+4
		•		ASEMBL	Integer+4
ITRN2	WELLSS	INTEGER*4		DUMP	INTEGER#4
				INITI	INTEGER*4
TRNDN	WELLSS	INTEGER*4		READ1	INTECER*4
ITRNI	ITER	Integer*4	ITSYX	HST3D	INTEGER*4
				ASEMBL	INTEGER*4
ITRNP	ITER	INTEGER*4		DUHP	INTEGER*4
				init1	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
ITSYZ	HST3D	INTEGER*4	ITWKT	HST3D	INTEGER*4
	ASEMBL	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INITI	INTEGER*4
	INITI	INTEGER*4		ITER	INTEGER*4
	READ1	Integer*4		READ1	INTEGER*4
	·_			SUMCAL	INTEGER*4
ITSZ	HST3D	INTEGER*4			
	ASEMBL	INTEGER*4	ITWSUR	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER*4
	INITI .	INTEGER*4		INITI	INTEGER*4
	READ1	INTEGER*4		ITER	INTEGER*4
				READ1	INTEGER*4
ITSZX	HST3D	INTEGER*4		SUMCAL	INTEGER*4
	Asembl	INTEGER*4			
	DUMP	INTEGER*4	ITX	HST3D	INTEGER*4
	INITI	integer*4		DUMP	INTEGER*4
	READ1	INTEGER*4		Initi	INTEGER*4
· .	•	1. <u>1</u> .		READ1	INTEGER*4
LTSZY	HST3D	INTEGER*4			
	Asembl	Integer*4	ITY	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER#4
	INITI	Integer*4		INITI	INTEGER*4
	READ1	INTEGER*4		READI	INTEGER#4
ITTWR	HST3D	INTEGER*4	ITZ	HST3D	INTEGER*4
	DUMP	INTEGER*4		DUMP	INTEGER#4
	INITI	INTEGER*4		INITI	INTEGER*4
	INIT2	INTEGER*4		READ1	INTEGER*4
	READ1	INTEGER*4			
	WBBAL .	INTEGER*4	IUBBLB	HST3D	INTEGER*4
	WELLSS	Integer*4			

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
IUCBC	HST3D	INTEGER*4	IVAIF	HST3D	INTEGER+4
				DUMP	Integer=4
IUDNBC	HST3D	INTEGER#4		initi	INTEGER#4
				READI	INTECER*4
Ludnlb	HST3D	Integer*4			
			IVASBC	HST3D	INTECER*4
LUDTHC	hst3d	INTEGER#4		DUMP	INTEGER#4
				INITI	INTEGER*4
LUH	D4DES	INTECER*4		iter	INTEGER#4
				READ1	Intecer=4
(UKHBC	HST3D	INTEGER#4			
			IVAW	hst3d	INTEGER#4
UKLB	HST3D	INTECER#4		DUMP	Integer*4
				initi	INTEGER#4
UPHIL	HST3D	INTEGER*4		ITER	INTECER*4
				READ1	INTEGER=4
UTBC	HST3D	Integer#4		SUNCAL	Intecer+4
UVAIF	HST3D	INTEGER#4	IVIS	hst3d	INTEGER±4
				DUAP	INTEGER#4
UVISL	HST3D	INTEGER*4		initi	INTEGER*4
				READ1	Intecer*4
IUZELB	HST3D	INTEGER#4	1		
			IVPA	hst3d	INTEGER*4 DIMENSION(*
EV	REWI3	INTEGER#4		DUMP	INTEGER*4 DIMENSION(*
				INIT1	INTEGER*4 DIMENSION(*
[VA	HST3D	INTEGER*4		ITER	INTEGER*4 DIMENSION(*
	DUMP	INTEGER*4		L2SOR	INTEGER*4 DIMENSION(*
	INIT1	INTEGER*4		READ1	INTEGER*4 DIMENSION(*
	L2SOR	INTEGER*4		SUHCAL	INTEGER*4 DIMENSION(*
	READ1	INTEGER*4			

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
IVSLBC	HST3D	INTEGER*4	IWEL	ASEMBL	INTEGER*4
	DUMP	INTEGER*4		ERROR2	INTEGER*4
	INITI	INTEGER*4		ERROR3	Integer*4
	READI	INTEGER*4		etom1	INTEGER*4
	•		•	INIT2	INTEGER#4
17XX	HST3D	INTEGER*4		ITER	INTEGER#4
,				PLOTOC	INTEGER#4
IVYY	HST3D	INTEGER*4		READ2	INTEGER*4
		- q.		READ3	INTEGER*4
IVZZ	HST3D	INTEGER*4		SUMCAL	INTEGER#4
		· ·		WBBAL	INTEGER*4
IW	ASEMBL	INTEGER*4 DIMENSION(*)		WELLSS	INTEGER#4
	ERROR2	INTEGER*4 DIMENSION(*)		WELRIS	INTEGER*4
	INIT2	INTEGER*4 DIMENSION(*)		WRITE2	INTEGER*4
	ITER	INTEGER*4 DIMENSION(*)		WRITE3	INTEGER*4
	READ2	INTEGER*4 DIMENSION(*)	•	WRITE5	Integer*4
	WEBAL	INTEGER*4 DIMENSION(*)			
	WELLSS	INTEGER*4 DIMENSION(*)	IWFICU	HST3D	Integer*4
	WRITE2	INTEGER*4 DIMENSION(*)		DUMP	INTEGER*4
	WRITE5	INTEGER*4 DIMENSION(*)		Init1	INTEGER*4
	ZONPLT	Integer*4		READ1	INTEGER*4
IWIP	WRITE5	INTEGER#4	IWFPCU	HST3D	INTEGER*4
				DUMP	INTEGER*4
IW2P	WRITE5	INTEGER*4		INIT1	INTEGER#4
	•			READ1	Integer*4
IWAIF	HST3D	INTEGER*4			
	DUMP	INTEGER*4	IWHICU	HST3D	Integer*4
	INITI	INTEGER*4		DUMP	INTEGER*4
	READ1	INTEGER*4		INIT1	INTEGER*4
				READ1	INTEGER*4
IWCF	HST3D	INTEGER*4			

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
IWHPCU	HST3D	INTEGER#4	I WQMTH	HST3D	INTEGER*4
	DUMP	INTEGER*4	•	DUMP	INTEGER*4
	initi	INTEGER*4		IN <b>ITI</b>	INTEGER*4
	READ1	INTEGER*4		ITER	INTEGER*4
				READ1	INTEGER*4
IWI	HST3D	INTEGER*4		SUHCAL	INTEGER*4
	DUMP	INTEGER*4			
	INITI	INTEGER*4	IWRANG	HST3D	INTEGER*4
	ITER	Integer#4		DUMP	INTEGER#4
	READ1	INTEGER#4		INITI	INTEGER*4
	SUMCAL	Integer#4		INIT2	INTEGER#4
				READ1	INTEGER*4
IWID	HST3D	INTEGER#4		WBBAL	INTEGER*4
	DUMP	INTECER#4		WELLSS	INTECER*4
	Initi	Intecer*4			
	INIT2	INTEGER=4	IWRSL	HST3D	INTEGER#4
	READ1	INTEGER*4		DUMP	INTEGER#4
	WBBAL	INTEGER*4		Init1	INTEGER#4
	WELLSS	INTEGER#4		INIT2	INTEGER*4
				READ1	INTEGER*4
IWOD	HST3D	INTEGER*4		WBBAL	INTEGER*4
	DUMP	INTEGER#4			
	INITI	INTEGER*4	IWRUF	HST3D	INTEGER*4
	INIT2	INTEGER*4		DUMP	INTEGER*4
	READ1	Integer*4		INIT1	INTEGER*4
	WBBAL	Integer*4		INIT2	INTEGER*4
	WELLSS	INTEGER*4		READ1	INTFGER*4
				WBBAL	INTEGER*4
IWPP	WRITES	INTEGER*4		WELLSS	INTEGER*4
IWQ1	WRITE2	INTEGER#4	IWSICU	HST3D	INTEGER*4
				DUMP -	INTECER*4
IWQ2	WRITE2	INTEGER*4		1NIT1	INTEGER*4
- •-				READ 1	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

	Refer-	•		Refer-	- · · · ·	
Variable	encing:	Variable	Variable	encing	Variable	
name	programs	type	name	programs	type	
IWSPCU	HST3D	INTEGER#4	IY	HST3D	INTEGER*4	
	DUMP	INTEGER*4		DUMP	INTEGER*4	
	INIT1	INTEGER#4		INITI	INTEGER*4	
	READ1	INTEGER*4		READI	INTEGER*4	
IX	HST3D	INTEGER*4	12	HST3D	INTEGER*4	
	DUMP	INTEGER*4		DUMP	INTEGER*4	
	INITI	INTEGER#4		INIT1	INTEGER*4	
	READ1	INTEGER#4		ITER	INTEGER*4	
IXIM	SOR2L	INTEGER*4		READ1	INTEGER*4	
IXIP	SOR2L	INTEGER#4	IZELBC	HST3D	INTEGER*4	
		*.		DUMP	INTEGER#4	
IX2M	SOR2L	Integer*4		INIT1	INTEGER*4	
				ITER	INTEGER*4	
1X2P	SOR2L	INTEGER#4		READ1	Integer*4	
IX3M	SOR2L	INTEGER*4	IZHCBC	HST3D	INTEGER*4	
		· · · · · · · · · · · · · · · · · · ·		DUMP	INTEGER*4	
IX3P	SOR2L	INTEGER*4		INIT1	INTEGER*4	
				READ1	INTEGER*4	
1XX	hst3d	INTEGER*4				
	DUMP	INTEGER*4			•	
	INITI	INTEGER*4			· · · · · · · · · · · · · · · · · · ·	
	L2SOR	Integer*4				
	READ1	INTEGER*4				
IXXN	HST3D	INTEGER*4			1.	
	DUMP	INTEGER*4				
-	INITI	INTEGER#4				
	L2SOR	INTEGER*4				
	READ1	INTEGER*4				

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
J	ASEMBL	INTEGER*4	JIZ	COEFF	INTEGER*4 DIMENSION(*)
	BSODE	INTEGER*4		ERROR2	INTEGER*4 DIMENSION(*)
	COEFF	INTEGER*4		INIT2	INTEGER*4 DIMENSION(*)
	CRSDSP	INTEGER+4		READ2	INTEGER*4 DIMENSION(*)
	D4DES	INTEGER=4		WRITE2	INTEGER*4 DIMENSION(*)
	DUMP	Integer#4			
	ERROR2	Intecer+4	J2	D4DES	INTEGER*4
	ETOMI	Intecer=4		IREWI	Integer#4
	INIT2	Integer+4		MAP2D	Integer+4
	INTERP	INTEGER#4		RFAD2	Integer+4
	IREWI	INTEGER*4		READ3	INTEGER*4
	MAP2D	Integer+4		REWI	INTEGER*4
	ORDER	INTEGER#4		REWI3	INTEGER+4
	READ1	INTEGER+4		ZONPLT	Integer#4
	READ2	INTECER+4			
	READ3	INTEGER#4	J2Z	COEFF	INTEGER#4 DIMENSION(*)
	REWI	INTEGER*4		ERROR2	INTEGER*4 DIMENSION(*)
	REWI3	INTEGER*4		INIT2	INTECER+4 DIMENSION (*)
	SUNCAL	INTEGER+4		READ2	INTEGER+4 DIMENSION(*)
	VSINIT	INTEGER*4		WRITE2	INTECER*4 DIMENSION(*)
	WRITES	INTEGER*4			
	ZONPLT	INTECER*4	J4	D4DES	INTEGER*4
<b>J1</b>	D4DES	INTECER#4	JBOT	MAP2D	INTEGER*4
	IREWI	Integer+4			
	MAP2D	Integer+4	JC	ERROR2	INTEGER*4
	READ2	Intecer+4		INIT2	INTEGER*4
	READ3	INTEGER*4		MAP2D	INTEGER*4
	REWI	INTECER*4			
	REWI3	INTECER*4	JCHAX	HST3D	INTEGER*4
	ZONPLT	Integer*4		SUMCAL	INTEGER*4
				WRITES	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
JHVSV	BSODE	INTEGER*4	JPRPTC	WRITE2 WRITE5	INTEGER*4 INTEGER*4
JHVSV1	BSODE	INTEGER*4		****	
JI	ERROR2	INTEGER*4	JPT	PLOTOC	INTEGER*4
			JSTART	BSODE	INTEGER*4
JIFMT	PRNTAR	INTEGER*4		WELRIS	INTEGER*4
JINC	MAP2D	INTEGER*4			
JING	TINE AU	ANADOMANY	JTIME	HST3D	INTEGER*4
JJ	D4DES	INTEGER#4		COEFF	INTEGER*4
J.J	DANCO	INIDUGATA	•	INIT3	INTEGER*4
JJ2	D4DES	INTEGER*4	JTHAX	HST3D	INTEGER*4
				SUMCAL	INTEGER*4
JL	PLOT	INTEGER*4		WRITE5	INTEGER*4
JMAP1	HST3D	INTEGER*4 DIMENSION(3)	JTOP	MAP2D	
	ERROR3				INTEGER*4
	ETOM2	INTEGER*4 DIMENSION(3)		• • • • • • • •	
	INIT3	INTEGER*4 DIMENSION(3)	JW	ASEMBL	INTEGER*4 DIMENSION(*)
	READ3	INTEGER*4 DIMENSION(3)		ERROR2	INTEGER*4 DIMENSION(*)
	WRITE3	INTEGER*4 DIMENSION(3)		INIT2	INTEGER*4 DIMENSION(*)
	WRITE5	INTEGER*4 DIMENSION(3)		ITER	INTEGER*4 DIMENSION(*)
	WALLGJ	ANALOLIN 4 DAILAIDADII(3)		READ2	INTEGER*4 DIMENSION(*)
JMAP2	HST3D	INTEGER*4 DIMENSION(3)	, .	WBBAL	INTEGER*4 DIMENSION(*)
	ERROR3	INTEGER*4 DIMENSION(3)		WELLSS	INTEGER*4 DIMENSION(*)
	ETOM2	INTEGER*4 DIMENSION(3)		WRITE2	INTEGER*4 DIMENSION(*)
	INIT3	INTEGER*4 DIMENSION(3)		WRITE5	INTEGER*4 DIMENSION(*)
	READ3	INTEGER*4 DIMENSION(3)			
	WRITE3	INTEGER*4 DIMENSION(3)	JWELL	WELLSS	INTEGER*4 DIMENSION(*)
	WRITES	INTEGER*4 DIMENSION(3)			
JODD	BSODE	INTEGER*4			
JPMAX	HST3D	INTEGER*4			
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Table 11.2--Cross-reference list of variables--Continued

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IPMAX HST3D INTEGER\*4 SUMCAL INTEGER\*4 WRITE5 INTEGER\*4

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
К	APLYBC	INTEGER#4	K1Z	COEFF	INTEGER*4 DIMENSION(*)
	ASEMBL	Integer+4		error2	INTEGER*4 DIMENSION(*)
	BSODE	INTEGER*4		INI <b>T</b> 2	INTEGER*4 DIMENSION(*)
	CALCC	Integer#4		READ2	INTEGER*4 DIMENSION(*)
	COEFF	INTEGER#4		WRITE2	INTEGER*4 DIMENSION(*)
	CRSDSP	INTEGER#4			
	DUMP	INTEGER#4	K2	IREWI	INTEGER*4
	ERROR2	INTEGER*4		MAP2D	INTEGER#4
	ETOM1	INTECER*4		REWI	INTEGER*4
	INIT2	INTEGER#4		REWI3	INTEGER*4
	IREWI	INTEGER#4			
	ITER	INTEGER*4	K2Z	COEFF	INTEGER*4 DIMENSION(*)
	MAP2D	INTEGER*4		error2	INTEGER*4 DIMENSION(*)
	ORDER	INTEGER*4		INIT2	INTEGER*4 DIMENSION(*)
	READ1	INTEGER*4		READ2	INTEGER*4 DIMENSION(*)
	READ2	INTEGER#4		WRITE2	INTEGER*4 DIMENSION(*)
	REWI	INTEGER*4			
	REWI3	INTEGER#4	KARHC	APLYBC	REAL*8 DIMENSION(*)
	SUMCAL	INTEGER*4	,	INIT2	REAL*8 DIMENSION(*)
	VSINIT	INTEGER*4			
	WBBAL	INTEGER*4	KC	PLOT	INTEGER*4
	WBCFLO	INTEGER*4			
	WELLSS	INTEGER*4	KCHAX	hst3d	INTECER*4
	WELRIS	1NTEGER*4		SUHCAL	INTEGER*4
	WRITE2	INTEGER*4		WRITE5	INTEGER*4
	WRITES	INTEGER*4			
			KF	INIT2	INTEGER*4
к1	INIT2	INTEGER*4			
	IREWI	INTEGER*4	KFLAG	BSODE	INTEGER*4
	MAP2D	INTEGER*4		WELRIS	INTEGER*4
	REWI	INTEGER*4			
	REWI3	INTEGER*4	KINC	init2	INTEGER#4
	WRITE5	INTEGER*4		,	
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		KK	MAP2D	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
KL	INIT2	INTEGER*4	KPMAX	HST3D	INTEGER*4
				SUMCAL	Integer*4
KLBC	APLYBC	REAL*8 DIMENSION(*)		WRITE5	INTEGER*4
	INIT2	REAL*8 DIMENSION(*)			
	WRITE2	REAL*8 DIMENSION(*)	KTHAWR	etom1	INTEGER*4 DIMENSION(*
		•		READ2	REAL*8 DIMENSION(*)
KMAP1	RST3D	INTEGER*4 DIMENSION(3)	÷	WELRIS	REAL*B
	ERROR3	INTEGER*4 DIMENSION(3)			
	ETOM2	INTEGER*4 DIMENSION(3)	KTHF	HST3D	REAL*8
	INIT3	INTEGER*4 DIMENSION(3)		DUMP	REAL*8
	READ3	INTEGER*4 DIMENSION(3)		ETOMI	REAL*8
	WRITE3	INTEGER*4 DIMENSION(3)		INIT2	REAL#8
	WRITE5	INTEGER*4 DIMENSION(3)		READ1	REAL*8
				READ2	REAL*8
KMAP2	RST3D	INTEGER#4 DIMENSION(3)		WRITE2	REAL <sup>48</sup>
	ERROR3	INTEGER*4 DIMENSION(3)			
	ETOM2	INTEGER*4 DIMENSION(3)	KTHWR	ETOM1	REAL*8 DIMENSION(*)
	INIT3	INTEGER*4 DIMENSION(3)		READ2	REAL*8 DIMENSION(*)
	READ3	INTEGER#4 DIMENSION(3)		WELRIS	REAL*8
	WRITE3	INTEGER*4 DIMENSION(3)			
	WRITE5	INTEGER*4 DIMENSION(3)	KTHX	COEFF	REAL*8 DIMENSION(*)
				etom1	REAL*8 DIMENSION(*)
KMAX	WELRIS	INTEGER*4		INIT2	REAL*8 DIMENSION(*)
ко	PLOT	INTEGER#4	KTHXPM	INIT2	REAL*8 DIMENSION(*)
				READ2	REAL*8 DIMENSION(*)
KOAR	HST3D	REAL*8		WRITE2	REAL*8 DIMENSION(*)
• .	APLYBC	REAL#8		•	
	DUMP	REAL*8	KTHY	COEFF	REAL*8 DIMENSION(*)
	ETOMI	REAL*8		ETOM1	REAL*8 DIMENSION(*)
ł	INIT2	REAL*8		INIT2	REAL*8 DIMENSION(*)
	READ1	REAL*8		•	
	READ2	REAL*8			
	WRITE2	REAL*8	}		

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
KTHYPM	INIT2	REAL*8 DIMENSION(*)	L	APLYBC	INTECER#4
	READ2	REAL*8 DIMENSION(*)		ASEMBL	INTEGER*4
	WRITE2	REAL*8 DIMENSION(*)		BSODE	INTEGER#4
				DUMP	INTECER#4
KTHZ	COEFF	REAL*8 DIMENSION(*)		Error2	INTECER#4
	ETOMI	REAL=8 DIMENSION(=)		ERROR3	INTEGER#4
	INIT2	REAL+8 DIMENSION(+)		INIT2	INTEGER#4
		•••		INIT3	INTEGER#4
KTHZPM	INIT2	REAL*8 DIMENSION(*)		ORDER	INTEGER*4
	READ2	REAL*8 DIMENSION(*)		PLOT	INTEGER#4
	WRITE2	REAL*8 DIMENSION(*)		READ1	INTEGER#4
				READ2	INTEGER#4
KTHAX	hst3d	INTEGER#4		SECFLO	Integer#4
	SUHCAL	INTEGER*4		SUHCAL	INTEGER*4
	WRITE5	INTEGER#4		WBCFLO	INTEGER#4
				WRITE2	INTEGER*4
KWEL	PLOTOC	INTEGER*4		WRITE3	INTEGER#4
				WRITES	INTEGER*4
KXX	ETOM1	REAL*8 DIMENSION(*)		ZONPLT	INTEGER#4
	init2	REAL*8 DIMENSION(*)			
	READ2	REAL*8 DIMENSION(*)	Ll	ERROR2	Integer=4
	WRITE2	REAL*8 DIMENSION(*)		INIT2	INTEGER#4
	•			ZONPLT	INTEGER#4
KYY	etom1	REAL*8 DIMENSION(*)			
	INIT2	REAL*8 DIMENSION(*)	L2	ERROR3	INTECER=4
	READ2	REAL*8 DIMENSION(*)		INIT2	Integer#4
	WRITE2	REAL*8 DIMENSION(*)		SOR2L	INTEGER#4
				ZONPLT	INTEGER*4
KZZ	eton1	REAL*8 DIMENSION(*)			
	INIT2	REAL*8 DIMENSION(*)	L3	D4DES	INTEGER#4
	READ2	REAL*8 DIMENSION(*)			
	WRITE2	REAL*8 DIMENSION(*)		•	

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
LABEL	REWI	CHARACTER*13 DIMENSION(5)	LCROSD	HST3D	LOGICAL*4
	REWI3	CHARACTER*10 DIMENSION(3)		ASEMBL	LOGICAL*4
	WRITE2	CHARACTER*20		COEFF	LOGICAL*4
	ZONPLT	CHARACTER*4		DUMP	LOGICAL*4
				ERROR1	LOGICAL*4
LBLDIR	SOR2L	CHARACTER*1 DIMENSION(3)		INIT1 ·	LOGICAL#4
	WRITE5	CHARACTER*1 DIMENSION(3)	•	INIT2	LOGICAL#4
				ITER	LOGICAL*4
LBLEQ	l2sor	CHARACTER*10 DIMENSION(3)		READ1	LOGICAL#4
	SOR2L	CHARACTER*10 DIMENSION(3)		READ2	LOGICAL#4
				SECFLO	LOGICAL#4
LBW	ORDER	INTEGER*4 DIMENSION(*)		WECFLO	LOGICAL*4
				WRITE1	LOGICAL#4
lcbotw	ASEMBL	INTEGER*4 DIMENSION(*)		WRITE2	LOGICAL*4
	Error2	INTEGER*4 DIMENSION(*)		WRITE4	LOGICAL#4
	INIT2	INTEGER*4 DIMENSION(*)		WRITES	LOGICAL#4
	ITER	INTEGER*4 DIMENSION(*)			
	READ2	INTEGER*4 DIMENSION(*)	LCTOPW	ASEMBL	INTEGER#4 DIMENSION(*)
	WBBAL	INTEGER*4 DIMENSION(*)		ERROR2	INTEGER#4 DIMENSION (*)
	WECFLO	INTEGER*4 DIMENSION(*)		INIT2	INTEGER*4 DIMENSION (*)
	WELLSS	INTEGER*4 DIMENSION(*)		ITER	INTEGER*4 DIMENSION (*)
	WRITE2	INTEGER*4 DIMENSION(*)		READ2	INTEGER*4 DIMENSION (*)
	WRITE5	INTEGER*4 DIMENSION(*)		WBBAL	INTEGER*4 DIMENSION(*)
				WBCFLO	INTEGER*4 DIMENSION (*)
				WELLSS	INTEGER*4 DIMENSION(*)
•				WRITE2	INTEGER*4 DIMENSION(*)
		··· ·· · · ·		WRITE5	INTEGER*4 DIMENSION (*)

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
			<u> </u>		
LDASH	WRITE5	CHARACTER*150	LINLIM	PLOT	INTEGER*4
LDOTS	CLOSE	CHARACTER*130	LL	APLYBC	INTEGER*4
	REWI	CHARACTER*130		INIT2	INTEGER#4
	REWI3	CHARACTER*120			
	WRITE2	CHARACTER#130	LLL	APLYBC	Integer+4
	WRITE3	CHARACTER#130			
	WRITES	CHARACTER*150	LN	MAP2D	INTEGER*4
LENAX	HST3D	REAL*8	LOCAIF	HST3D	INTEGER*4
	MAP2D	REAL*8		APLYBC	INTEGER*4
	READ3	REAL+8		DUMP	INTEGER*4
	WRITE3	REAL#8		ERROR2	INTEGER*4
	WRITES	REAL*8		ETOM1	INTEGER*4
	ZONPLT	REAL*8		INIT2	INTEGER*4
				READI	INTEGER*4
LENAY	HST3D	REAL*8		READ2	INTEGER*4
	HAP2D	REAL*8		WRITE2	INTEGER#4
	READ3	REAL*8			
	WRITE3	REAL*8	LPRNT	PRNTAR	INTEGER*4 DIMENSION(*)
	WRITES	REAL#8		WRITE2	INTEGER*4 DIMENSION(*)
	WALLEJ			WRITE3	INTEGER*4 DIMENSION(*)
LENAZ	HST3D	REAL*8		WRITE4	INTEGER*4 DIMENSION(*)
LERAL	READ3	REAL#8		WRITE5	INTEGER*4 DIMENSION(*)
				*****	
	WRITE3	REAL#8	LRBC	ETOM1	INTEGER*4
	WRITE5	REAL*8		~4 \415	L # L # J # K ** 7
LGREN	WELLSS	REAL*8	LTD	APLYBC	REAL*8
	WFDYDZ	REAL*8	_		
			LXL	HAP2D	INTEGER*4
LIMAGE	READ1	CHARACTER*80			
LIMIT	WRITE3	CHARACTER*4			

Table 11.2--Cross-reference list of variables--Continued

PLOT

LINE

CHARACTER\*101

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
LXR	MAP2D	INTEGER*4	Ml	ORDER	INTEGER*4
			•••	REWI	INTEGER*4
M	APLYBC	INTEGER*4		REWI 3	INTEGER*4
••	ASEMBL	INTEGER*4		SOR2L	INTEGER*4
	BSODE	INTEGER*4		WELLSS	INTEGER*4
	COEFF	INTEGER*4			
	CRSDSP	INTEGER*4	M2	BSODE	INTEGER#4
	D4DES	INTEGER*4		REWI	INTEGER*4
	DUMP	INTEGER*4		REWI3	INTEGER*4
	ERROR2	INTEGER#4		SOR2L	INTEGER*4
	ERROR3	INTEGER*4			· · · · · · · · · · · · · · · · · · ·
	ETOM1	INTEGER*4	M3	SUMCAL	INTEGER#4
	ETOM2	INTEGER*4			
	INIT2	INTEGER*4	MA	ASEMBL	INTEGER*4
	INIT3	INTEGER*4		SBCFLO	INTEGER*4
	IREWI	INTEGER*4		WBCFLO	INTEGER*4
	ITER	INTEGER#4			
	L2SOR	INTEGER*4	MAIFC	APLYBC	INTEGER*4 DIMENSION(*)
	ORDER	INTEGER*4		ASEMBL	INTEGER#4 DIMENSION(#
	PRNTAR	INTEGER*4		ERROR3	INTEGER*4 DIMENSION(*)
	READ1	INTEGER*4		INIT2	INTEGER*4 DIMENSION(*)
	READ2	INTEGER*4		INIT3	INTEGER*4 DIMENSION(*)
	READ3	INTEGER#4		SUMCAL	INTEGER*4 DIMENSION(*)
	REWI	INTEGER*4		WRITE2	INTEGER*4 DIMENSION(*)
	REWI3	INTEGER#4		WRITE5	INTEGER*4 DIMENSION(*)
•	SBCFLO	INTEGER*4		•	
	SOR2L	INTEGER#4	MAPPTC	HST3D	INTEGER*4
	SUMCAL	INTEGER*4		ERROR3	INTEGER*4
•	WBBAL	INTEGER#4		READ3	INTEGER#4
-	WBCFLO	INTEGER*4		WRITE3	INTEGER*4
	WELLSS	INTEGER*4		WRITE5	INTEGER*4
	WRITE2	INTEGER*4		•	
	WRITE3	INTEGER*4	MAXDXX	SOR2L	REAL*8
	WRITE4	INTEGER*4			
	WRITE5	INTEGER*4	MAXIT	SOR2L	INTEGER*4

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
MAXITI	HST3D	INTEGER*4	METH	HST3D	INTEGER*4
	DUMP	INTEGER*4		BSODE	INTEGER*4
	INIT2	INTEGER*4		WELRIS	INTEGER*4
	READ1	INTEGER*4			
	READ2	INTEGER*4	MFBC	APLYBC	INTEGER*4 DIMENSION(*)
	SOR2L	INTEGER*4		ASEMBL	INTEGER#4 DIMENSION (#)
			•	ERROR3	INTEGER*4 DIMENSION(*)
MAXIT2	hs <b>t</b> 3d	INTEGER*4		INIT2	INTEGER#4 DIMENSION(*)
	DUMP	Intecer+4		INIT3	INTEGER*4 DIMENSION(*)
	INIT2	Integer#4		SUMCAL	INTEGER*4 DIMENSION(*)
	READ1	Integer+4		WRITE2	INTEGER*4 DIMENSION(*)
	READ2	Intecer+4		WRITE3	INTEGER*4 DIMENSION(*)
	SOR2L	INTECER*4		WRITES	INTECER*4 DIMENSION(*)
MAXITN	HST3D	INTEGER*4	MFLBL	PLOTOC	CHARACTER*12
	DUMP	Intecer+4		WRITE2	CHARACTER*12
	INIT2	Intecer*4		WRITE3	CHARACTER*12
	ITER	Integer+4		WRITE5	CHARACTER*12
	READ1	INTECER*4			
	READ2	INTEGER*4	MHCBC	APLYBC	INTEGER*4 DIMENSION(*)
	WRITE2	INTEGER*4		ASEMBL	INTEGER*4 DIMENSION(*)
				INIT2	INTEGER*4 DIMENSION(*)
MAXORD	HS <b>T3D</b>	INTEGER*4		SUHCAL	INTEGER*4 DIMENSION(*)
	BSODE	INTEGER*4		WRITE2	INTEGER*4 DIMENSION(*)
	WELRIS	INTECER*4		WRITE5	INTECER*4 DIMENSION(*)
MAXPTS	HST3D	INTEGER*4	MIJKM	HST3D	INTEGER*4
	BSODE	INTEGER*4		ASEMBL	INTEGER*4
	WELRIS	INTEGER*4		COEFF	INTEGER*4
MAXXX	SOR2L	REAL*8		CRSDSP	INTEGER*4
				INIT2	INTEGER*4
HD4	D4DES	INTEGER*4		SBCFLO	INTEGER*4
	ORDER	INTEGER*4		WBBAL	INTEGER*4
				WBCFLO	INTEGER*4
				WELLSS	1NTEGER*4

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
MIJKP	HST3D	INTEGER*4	MIMJK	HST3D	INTEGER*4
	ASEMBL	INTEGER*4		ASEMBL	· INTEGER#4
	COEFF	Integer*4		CRSDSP	INTEGER*4
	CRSDSP	INTEGER*4		SBCFLO	Integer*4
	SBCFLO	INTEGER*4		WECFLO	Integer*4
	WBBAL	INTEGER*4	· ·		
	WECFLO	INTEGER#4	MIMJKM	CRSDSP	INTEGER*4
	WELLSS	INTEGER*4			
			MIMJKP	CRSDSP	Integer*4
MIJMK	HST3D	INTEGER*4			
	ASEMBL	INTEGER*4	MIMJMK	CRSDSP	INTEGER*4
	COEFF	INTEGER*4			
	CRSDSP	INTEGER#4	MIMJPK	CRSDSP	INTEGER*4
•	INIT2	INTEGER#4			
	SBCFLO	INTEGER*4	MINUS	PLOT	CHARACTER*1
	WBCFLO	INTEGER*4			-
			MIPJK	HST3D	Integer*4
MIJMKH	CRSDSP	INTEGER*4	•	ASEMBL	INTEGER*4
				CRSDSP	INTEGER*4
MIJMKP	CRSDSP	INTEGER*4		SBCFLO	INTEGER*4
	. ·			WECFLO	INTEGER*4
MIJPK	HST3D	INTEGER*4			
	ASEMBL	INTEGER*4	MIPJKM	CRSDSP	Integer*4
	COEFF	INTEGER*4			
	CRSDSP	INTEGER*4	MIPJKP	CRSDSP	INTEGER*4
	SBCFLO	INTEGER*4			
	WECFLO	INTEGER*4	MIPJMK	CRSDSP	INTEGER*4
MIJPKM	CRSDSP	INTEGER*4	MIPJPK	CRSDSP	INTEGER*4
MIJPKP	CRSDSP	INTEGER*4			

	Refer-			Refer-	
Variable	encing	Variable	Variable	~	Variable
name	programs	type	name	programs	type
мк	INIT2	INTEGER*4	нт	COEFF	INTEGER*4
				INIT2	INTEGER=4
MKT	WBBAL	INTEGER*4		SUNCAL	INTEGER*4
	WELLSS	INTEGER*4			
	WRITES	INTEGER#4	MTJP1	COEFF	INTEGER*4
HL	INIT2	INTEGER*4	htwo	BSODE	INTEGER*4
MLBC	APLYBC	INTECER*4 DIMEN		INIT2	INTEGER*4
	ASEMBL	INTEGER*4 DIMEN			
	ERROR2	INTEGER*4 DIMEN		HST3D	INTEGER*4
	INIT2	INTEGER*4 DIMEN		DUMP	INTEGER*4
	INIT3	INTEGER*4 DIMEN		etom1	INTECER*4
	SUNCAL	INTEGER#4 DIMEN	• •	init2	INTEGER*4
	WRITE2	INTEGER*4 DIMEN		READ1	INTEGER*4
	WRITE3	INTECER#4 DIMEN		READ2	INTEGER*4
	WRITE5	INTEGER*4 DIMEN	SION(*)	WELLSS	INTEGER#4
				WRITE2	INTEGER*4
MM	ASEMBL	INTEGER*4			
	INIT2	INTECER*4 DIMEN	5ION(8) N	MAP2D	INTEGER*4
	ITER	INTEGER#4			
	WBBAL	Integer+4	N1	PRNTAR	INTEGER*4
	WELLSS	INTEGER*4		ZONPLT	INTEGER*4
	<b>WRITES</b>	INTECER*4			
			N2	PRNTAR	INTEGER*4
MNEXT	BSODE	INTECER*4		ZONPLT	INTEGER*4
MOBW	WELLSS	REAL*8 DIMENSIO	N(*) N3	PRNTAR	INTEGER*4
				ZONPLT	INTEGER*4
MSBC	ASEMBL	INTEGER*4 DIMEN	• •		
	INIT2	INTEGER*4 DIMEN	• -	MAP2D	INTEGER*4
	INIT3	INTEGER*4 DIMEN			
	SBCFLO	INTECER*4 DIMEN		PRNTAR	CHARACTER*6
	SUHCAL	INTEGER*4 DIMEN			
	WRITE2	INTEGER*4 DIMEN		PRNTAR	CHARACTER*2
	WRITE3	INTEGER*4 DIMEN			
	WRITE5	INTEGER*4 DIMEN	SION(*) NAC	ERROR2	INTEGER*4

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	' Variable type
NAIFC	HST3D	INTEGER*4	NDELA	MAP2D	INTEGER*4
	APLYBC	INTEGER*4			
	ASEHBL	INTEGER*4	NDIM	INTERP	INTEGER*4
•	DUMP	INTEGER*4		PRNTAR	INTEGER*4
	ERRORI	INTEGER*4			
	ERROR2	INTEGER*4	NEHST	HST3D	INTEGER*4
	ERROR3	INTEGER#4			INTEGER*4
	ETOM1	INTEGER*4		DUMP	INTEGER*4
	ETOM2	INTEGER*4		INIT2	INTEGER*4
	INITI	INTEGER*4		READI	INTEGER*4
	INIT2	INTEGER*4		TOFEP '	INTEGER#4
INI Ite	INIT3	INTEGER*4			
	ITER	INTEGER*4	NFBC	HST3D	INTEGER*4
	READ1	INTEGER*4		APLYBC	INTEGER*4
	READ2	INTEGER*4		ASEMBL	INTEGER*4
	READ3	INTEGER*4		DUMP	INTEGER*4
	SUMCAL	INTEGER*4		ERROR1	INTEGER*4
	WRITE1	INTEGER*4		ERROR2	INTEGER*4
	WRITE2	INTEGER*4		ERROR3	INTEGER*4
	WRITE3	INTEGER*4		ETOMI	INTEGER*4
	WRITE5	INTEGER*4		ETOM2	INTEGER*4
				INITI	INTEGER*4
NC	PLOT	INTEGER*4		INIT2	Integer*4
	PLOTOC	INTEGER*4		INIT3	INTEGER*4
		• •	,	ITER	INTEGER*4
NCHARS	MAP2D	INTEGER*4	·	READ1	INTEGER*4
				READ2	INTEGER*4
NCHP	MAP2D	INTEGER*4		READ3	INTEGER*4
	ZONPLT	INTEGER*4		SUMCAL	INTEGER*4
		•		WRITEI	INTEGER*4
NCHPL	MAP2D	INTEGER*4		WRITE2	INTEGER*4
				WRITE3	INTEGER*4
NCHPR	MAP2D	INTEGER*4		WRITE5	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Vari: tyj		Variable name	Refer- encing programs	Variable type
NGRIDX	MAP2D	INTEGER*4	DIMENSION (50)	NHCN	HST3D	INTEGER*4
					APLYBC	INTEGER*4
NGRIDY	MAP2D	INTEGER*4	DIMENSION(50)		DUMP	INTEGER+4
					ERROR1	INTEGER*4
NHC	ERROR2	1ntecer+4			ERROR2	Integer*4
					etom1	INTEGER#4
NHCBC	HST3D	INTEGER*4			INITI	INTECER*4
	APLYBC	INTEGER#4			INIT2	INTEGER*4
	ASEMBL	INTEGER*4			READ1	INTEGER#4
	DUMP	INTEGER#4			READ2	Integer*4
	ERROR2	INTEGER#4			WRITE1	INTEGER*4
	ERROR3 ETOM1	INTECER*4 INTECER*4			WRIŢE2	Integer*4
	ETOM1 ETOM2	INTEGER*4		1.TT		Thursday and I
	INITI	INTEGER#4		NL	ZONPLT	INTEGER*4
	INIT2	INTEGER#4			•	
	INIT3	INTEGER*4				
	ITER	INTEGER#4				
	READ1	INTEGER*4				
	READ2	INTEGER*4				
	READ3	INTEGER*4				
	SUMCAL	INTEGER*4				
	WRITE1	INTEGER*4				
	WRITE2	INTEGER*4				
	WRITE3	INTECER*4				
	WRITES	INTEGER*4				

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
NLBC	HST3D	INTEGER*4	NMPZON	HST3D	INTEGER*4 DIMENSION(3)
	APLYBC	INTEGER*4		INIT3	INTEGER*4 DIMENSION(3)
•	ASEMBL	INTEGER*4		MAP2D	INTEGER*4
	DUMP	INTEGER*4		READ3	INTEGER*4 DIMENSION(3)
:	ERROR1	INTÉGER*4		WRITE3	INTEGER*4 DIMENSION(3)
	ERROR2	INTEGER*4		WRITE5	INTEGER*4 DIMENSION(3)
	ERROR3	INTEGÉR*4	•••		
	etom1	Integer#4	NN3	ZONPLT	INTEGER*4
	ETOM2	INTEGER*4			
	INITI	INTEGER*4	NNC	D4DES	INTEGER*4
	INIT2	INTEGER*4			
	INIT3	INTEGER#4	NNOPPR	PRNTAR	INTEGER*4
	ITER	INTEGER*4	÷		
	READ1	INTEGER*4	NNOUT	MAP2D	INTEGER*4
	READ2	INTEGER*4			
	READ3	INTEGER*4	NNPR	PRNTAR	INTEGER*4
n a na	SUMCAL	INTEGER*4			
	WRITE1	INTEGER*4	NO	PLOT	INTEGER*4
	WRITE2	Integer*4		PLOTOC	INTEGER*4
	WRITE3	INTEGER*4		4	
	WRITE5	INTEGER*4	NOCV	HST3D	INTEGER*4
		•		DUMP	INTEGER*4
NLC	ERROR2	INTEGER#4		ERROR2	INTEGER*4
	11 - 11 - 11 - 11 - 11 - 11 - 11 - 11	÷ .		ETOMI	INTEGER*4
NLP	PLOT	INTEGER*4		INIT2	INTEGER*4
	ZONPLT	INTEGER*4		READ1	INTEGER*4
		•		READ2	INTEGER*4
NMAPR	HST3D	INTEGER*4		VISCOS	INTEGER*4
:	CLOSE	INTEGER*4		WRITE2	INTEGER*4
·· · .	DUMP	INTEGER*4			
	INIT2	INTEGER*4	NOTV	VSINIT	INTEGER*4
	READ1	INTEGER*4			

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	Refer-			Refer-	
Variable	encing	Variable	Variable	encing	Variable
name	programs	type	name	programs	type
NOTVO	HST3D	INTEGER*4	NPMZ	HST3D	INTEGER#4
	DUMP	INTECER*4		COEFF	INTEGER*4
	ERROR2	INTEGER*4		DUMP	INTEGER#4
	etom1	INTEGER*4		ERROR 1	INTEGER*4
'	INIT2	INTEGER*4		ERROR2	INTEGER*4
	READ1	INTEGER*4		etom1	INTEGER#4
	READ2	INTEGER*4		INITI	<b>1NTEGER*4</b>
	VISCOS	INTEGER*4		INIT2	INTEGER#4
	WRITE2	INTEGER*4		READ1	INTECER#4
				kEAD2	INTEGER*4
Notv1	hst3d	INTECER*4		WRITEL	INTEGER*4
	DUMP	INTEGER*4		WRITE2	INTEGER#4
	ERROR2	INTEGER*4		ZONPLT	INTEGER*4
	etom1	INTEGER#4			
	INIT2	INTEGER*4	nposns	HAP2D	INTEGER*4
	READI	INTEGER#4			
	READ2	INTEGER*4	NPR2	PRNTAR	INTEGER*4
	VISCOS	INTEGER*4			
	WRITE2	INTECER*4	NPR3	PRNTAR	INTEGER*4
NP	ZONPLT	INTEGER#4	NPTAIF	HST3D	INTEGER#4
				DUMP	INTEGER*4
NPAGES	HAP2D	INTEGER*4		ERROR2	INTEGER#4
				ETOHI	INTEGER#4
NPAIF	APLYBC	INTEGER*4		INIT2	INTEGER*4
		••		READ1	INTEGER#4
NPCX	ZONPLT	INTEGER*4		READ2	INTEGER*4
				WRITE2	INTEGER*4
NPCY	ZONPLT	1NTEGER*4			
NPEHDT	HST3D	INTEGER*4			
-	<b>BLOCKDATA</b>	INTEGER*4			
	DUHP	INTEGER*4			
	INIT2	INTEGER*4			
	READ1	INTEGER*4			

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Variable	Refer- encing	Variable	Variable	Refer- encing	Variable
name	programs	type	name	programs	type
NPTCBC	HST3D	INTEGER*4	NPTSD4	HST3D	INTEGER*4
	APLYBC	INTEGER*4		D4DES	INTEGER*4
	ASEMBL	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		INITI	Integer*4
	ERRORI	INTEGER*4		ORDER	INTEGER*4
	ERROR2	INTEGER*4	1	READI	INTEGER*4
	ERROR3	INTEGER*4		WRITE1	INTEGER*4
- 1	ETOMI	INTEGER*4			
	ETOM2	INTEGER#4	NPTSUH	HST3D	INTEGER*4
	INITI	INTEGER*4		ASEMBL	Integer*4
	INIT2	INTEGER*4		D4DES	INTEGER*4
	INIT3	INTEGER*4		DUMP	Integer*4
	ITER	INTEGER*4		init1	INTEGER*4
	READ1	INTEGER*4		ORDER	INTEGER*4
	READ2	INTEGER#4		READI	INTEGER*4
	READ3	INTEGER*4		SBCFLO	INTEGER*4
	SBCFLO	INTEGER*4		WBCFLO	INTEGER*4
	SUMCAL	INTEGER*4		WRITE1	INTEGER*4
	WRITEI	INTEGER*4			
	WRITE2	INTEGER#4	NR	ERROR2	INTEGER*4
	WRITE3	INTEGER*4		INIT2	Integer*4
	WRITE5	INTEGER#4		READ2	INTEGER*4
	•			WRITE2	Integer#4
NPTSA4	HST3D	INTEGER*4			
•••• • • • • • • • •	DADES	INTEGER*4	NROWS	MAP2D	INTEGER*4
	DUMP	INTEGER*4			
	INITI	INTEGER*4	NRP	ZONPLT	Integer*4
•	ORDER	INTEGER*4		. •	
	READI	INTEGER*4	NRPR	ZONPLT	INTEGER*4
	WRITE1	INTEGER*4			

Table 11.2--Cross-reference list of variables--Continued

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/ariable name	Refer- encing programs	Variable Lype	Variable name	Refer- encing programs	Variable type
NRSTTP	HST3D	INTEGER*4	NTHPTO	PLOTOC	INTEGER*4
	CLOSE	INTEGER*4			
	DUHP	INTEGER*4	NTSCHK	HST3D	INTEGER*4
	INLT2	INTEGER=4		CLOSE	INTEGER#4
	READ1	Intecer+4		Dump	Integer+4
				READ3	INTEGER*4
NS	ZONPLT	INTECER#4			
			NTSOPT	HST3D	INTEGER*4
NSC	ERROR2	INTEGER#4		DUMP	INTEGER#4
				INIT2	INTEGER*4
NSHUT	WELLSS	INTEGER*4		ITER	INTEGER*4
				READ1	INTEGER*4
nstd4	HST3D	INTEGER*4		READ2	INTEGER#4
	DUMP	INTEGER*4		SOR2L	INTEGER*4
	INITI	INTEGER*4		WRITE2	INTEGER#4
	READ1	INTEGER*4		~~~~~~	
	WRITEI	INTEGER*4	NVST	VSINIT	INTECER+4
	WALLUL			TULMEL	
NSTSOR	HST3D	INTEGER*4			
	DUMP	INTECER+4			
	INITI	INTEGER#4			
	READ1	INTEGER*4			
	WRITE1	INTEGER*4			
NTEHDT	HST3D	INTEGER#4			
		INTEGER*4			
	DUMP	INTEGER*4			
	INIT2	INTEGER*4			
	READ1	INTEGER*4			
	TOFEP	INTEGER*4			
NTHPTC	PLOTOC	1NTEGER*4			

Table	11.2Cross-reference	list of	variablesContinued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
WEL	HST3D	INTEGER*4	NX	HST3D	INTEGER*4
	ASEMBL	Integer#4		APLYBC	INTEGER*4
	DUMP	INTEGER*4		ASEMBL	INTEGER*4
	ERROR2	Integer*4		COEFF	INTEGER*4
	ERROR3	INTEGER*4		CRSDSP	Integer*4
	etom1	Integer*4		D4DES	INTEGER*4
	ETOM2	Integer*4		DUMP	INTEGER*4
	initi	Integer*4		ERROR1	INTEGER*4
	INIT2	INTEGER*4		ERROR2	Integer*4
	INIT3	INTEGER*4		ETOMI	INTEGER*4
	ITER	INTEGER*4		etom2	Integer#4
	READ1	Integer*4		INITI	INTEGER*4
	READ2	INTEGER*4		INIT2	INTEGER*4
	READ3	Integer*4		INIT3	INTEGER*4
	SUMCAL	Integer*4		INTERP	INTEGER*4
	WBBAL	INTEGER*4		IREWI	INTEGER*4
	WELLSS	INTEGER*4		ITER	INTEGER*4
	WRITE1	INTEGER*4		MAP2D	INTEGER*4
	WRITE2	INTEGER*4		ORDER	INTEGER*4
	WRITE3	INTEGER*4		PLOTOC	INTEGER#4
	WRITE5	INTEGER*4		PRNTAR	INTEGER*4
				READI	INTEGER*4
				READ2	INTEGER*4
				READ3	Integer*4
	2 .	and the second		REWI	INTEGER*4
				REWI3	INTEGER*4
				SECFLO	INTEGER*4
		-		SOR2L	INTEGER*4
				SUMCAL	INTEGER*4
		· · ·		WBBAL	INTEGER*4
				WBCFLO	INTEGER*4
		5		WELLSS	INTEGER*4
				WRITE1	INTEGER*4
				WRITE2	INTEGER*4
				WRITE3	INTEGER*4
				WRITE4	INTEGER*4
				WRITE5	INTEGER*4
				ZONPLT	INTEGER*4

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Variable name	Refer- encing programs	Variable type	Variable . name	Refer- encing programs	Variable type
NX 1	ORDER	INTEGER#4	NXY	HST3D	INTEGER*4
	SOR2L	INTEGER*4		APLYBC	INTEGER*4
	ZONPLT	INTEGER*4		ASEMBL	INTEGER*4
				COEFF	INTECER*4
NX2	ORDER	Integer+4		CRSDSP	INTEGER*4
	SOR2L	Integer*4		D4DES	INTECER*4
	ZONPLT	Integer#4		DUMP	INTEGER*4
				Error1	INTECER*4
NX3	ORDER	Intecer=4		ERROR2	INTEGER*4
	SOR2L	Integer+4		etom1	INTEGER*4
				ETOM2	INTEGER*4
IXPR	PRNTAR	INTEGER#4		iņiti	INTEGER*4
				INIT2	INTECER*4
XX	INIT2	integer*4		INIT3	INTEGER#4
	TOFEP	INTECER*4		IREWI	INTEGER*4
	VISCOS	INTEGER#4		ITER	INTEGER#4
				ORDER	INTECER*4
				PLOTOC	Integer+4
				PRNTAR	INTEGER*4
				READ1	INTECER*4
				READ2	INTECER*4
				READ3	INTEGER*4
				REWI	INTEGER*4
				REWI3	INTECER*4
				SBCPLO	INTEGER*4
				SOR2L	Integer*4
				SUMCAL	INTEGER#4
				WBBAL	Integer*4
				WBCFLO	INTEGER#4
				WELLSS	Integer*4
				WRITE1	INTEGER*4
				WRITE2	INTEGER*4
				WRITE3	INTEGER*4
				WRITE4	INTEGER*4
				WRITE5	INTEGER*4
			(		

Table 11.2--Cross-reference list of variables--Continued

	Refer-			Refer-	· · · · · · · · · · · · · · · · · · ·
Variable name	encing programs	Variable type	Variable name	encing programs	Variable type
110111C	Programs				
NXYZ	HST3D <sup>®</sup>	INTEGER*4	NY	HST3D	INTEGER*4
	APLYBC	INTEGER*4		APLYBC	INTEGER*4
	ASEMBL	Integer*4		ASEMBL	1nteger*4
	COEFF	INTEGER*4		COEFF	INTEGER*4
	D4DES	INTEGER*4		DUMP	INTEGER*4
	DUMP	INTEGER*4		ERRORI	INTEGER*4
	ERROR1	INTÉGER#4		ERROR2	INTEGER*4
	ERROR2	INTEGER*4		ETOMI	INTEGER*4
	ETOM1	Integer#4		INIT1	INTEGER*4
	ETOM2	INTEGER*4		INIT2	INTEGER*4
	INITI	INTEGER*4		INIT3	INTEGER*4
	INIT2	INTEGER*4		INTERP	INTEGER*4
	INIT3	INTEGER*4		IREWI	INTEGER*4
	IREWI	INTEGER*4		ITER	1NTEGER*4
	ITER	INTEGER*4		MAP2D	INTEGER*4
	LZSOR	INTEGER*4		ORDER	INTEGER*4
	ORDER	INTEGER*4		PRNTAR	INTEGER#4
	PLOTOC	INTEGER*4		READ1	INTEGER*4
	PRNTAR	INTEGER*4		READ2	INTEGER*4
	READ1	INTEGER*4		READ3	INTEGER*4
	READ2	INTEGER*4		REWI	INTEGER*4
	READ3	INTEGER*4		REWI3	INTEGER*4
	REWI	INTEGER*4		SOR2L	INTEGER#4
	REWI3	INTEGER*4		SUMCAL	INTEGER#4
	SOR2L	INTEGER*4	1	WBBAL	INTEGER*4
	SUMCAL	1NTEGER*4		WELLSS	INTEGER*4
	WBBAL	INTEGER*4	•	WRITEL	INTEGER*4
	WELLSS	INTEGER*4		WRITE2	INTEGER*4
	WRITE!	1NTEGER*4		WRITE5	INTEGER*4
	WRITE2	INTEGER*4	·	ZONPLT	INTEGER*4
	WRITE3	INTEGER*4		· ·	
	WRITE4	INTEGER*4	NYI	ZONPLT	INTEGER*4
	WRITE5	INTEGER*4		•	
			NY 2	ZONPLT	INTEGER*4

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
NYPR	PENTAR	INTEGER*4	NZTPHC	HST3D	INTEGER*4
				DUMP	INTEGER#4
NZ	HST3D	INTEGER*4		ETOH1	INTEGER#4
	APLYBC	INTEGER*4		INIT2	INTEGER*4
	ASEMBL	INTECER=4		READ1	INTEGER*4
	CALCC	INTEGER#4		READ2	INTEGER*4
	COEFF	INTECER*4		WRITE2	INTEGER*4
	DUMP	INTEGER*4			
	ERROR1	INTEGER#4	NZTPRO	HST3D	INTEGER*4
	ERROR2	Integer#4		DUMP	INTEGER*4
	ETOM1	INTEGER#4		etomi	INTEGER*4
	INIT1	INTEGER#4		INIT2	INTEGER#4
	INIT2	INTEGER#4		READ1	INTEGER*4
	INIT3	INTEGER#4		READ2	INTEGER*4
	IREWI	INTEGER*4		WRITE2	INTEGER*4
	ITER	INTEGER#4			
	ORDER	INTEGER#4	OCPLOT	HST3D	LOGICAL*4
	PRNTAR	INTEGER*4		CLOSE	LOGICAL*4
	READI	INTEGER#4		DUMP	LOGICAL*4
	READ2	INTEGER*4		ERROR3	LOGICAL*4
	READ3	INTEGER*4		READ1	LOGICAL*4
	REWI	INTEGER*4		READ2	LOGICAL*4
	REWI3	INTEGER*4		READ3	LOGICAL*4
	SOR2L	INTEGER*4		SOR2L	LOGICAL*4
	SUMCAL	INTEGER*4		WELLSS	LOGICAL*4
	WBBAL	INTEGER*4		Welris	LOGICAL*4
	WELLSS	1NTEGER*4		WRITE2	LOGICAL*4
	WRITE1	INTEGER*4		WRITE3	LOGICAL*4
	WRITE2	INTEGER*4		WRITE5	LOGICAL*4
	WRITE5	INTEGER*4			
			ODD	ORDER	LOGICAL*4
NZPR	PRNTAR	INTEGER*4	<b></b>		•••••
			он	PLOT	CHARACTER*1

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
OMEGA	L2SOR	REAL*8	Р	APLYBC	REAL*8 DIMENSION(*)
	SOR2L	REAL*8		ASEMBL	REAL*8 DIMENSION(*)
				CALCC	REAL*8
OMGMAX	SOR2L	REAL*8		COEFF	REAL*8 DIMENSION(*)
		÷		ETOML	REAL*8 DIMENSION(*)
OMGMIN	SOR2L	REAL*8		INIT2	REAL*8 DIMENSION(*)
				ITER	REAL*8 DIMENSION(*)
OMOPT	HST3D	REAL*8 DIMENSION(3)		READ2	REAL*8 DIMENSION(*)
	DUMP	REAL*8 DIMENSION(3)		SUMCAL	REAL*8 DIMENSION(*)
	L2SOR	REAL*8 DIMENSION(3)		TOFEP	REAL*8
	READ1	REAL*8 DIMENSION(3)		WBBAL	REAL*8 DIMENSION(*)
				WELLSS	REAL*8 DIMENSION(*)
ORENPR	HST3D	INTEGER*4		WRITE2	REAL*8 DIMENSION(*)
	DUMP	INTEGER*4		WRITE5	REAL*8 DIMENSION(*)
	INIT2	INTEGER#4		ZONPLT	CHARACTER*10000
	PRNTAR	INTEGER*4		10 A	
	READI	INTEGER#4	PO	HST3D	REAL*8
	READ2	INTEGER*4		DUMP	REAL*8
	WRITE2	INTEGER*4		ETOM1	REAL*8
	WRITE3	INTEGER*4		INIT2	REAL*8
	WRITE4	INTEGER*4		READ1	REAL*8
	WRITE5	INTEGER*4		READ2	REAL*8
	•			SUMCAL	REAL*8
ORFR	WRITE5	INTEGER*4		WBBAL	REAL*8
		••••••		WELLSS	REAL*8
		_ · · · ·		WPDYDZ	REAL*8
		. · ·		WRITE2	REAL*8
	-			WRITE5	REAL*8

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variabl <i>e</i> type	Variable name	Refer- encing programs	Variable type
P00	HST3D	REAL*8	PAATM	HST3D	REAL*8
	WBBAL	REAL*8		Аргувс	REAL*8
	WELLSS	REAL*8		ASEMBL	REAL*8
	WELRIS	REAL*8		DUMP	REAL*8
				etohi	REAL*8
рон	HST3D	REAL#8		INIT2	REAL*8
	APLYBC	REAL*8		READ1	REAL*8
	ASEMBL	REAL*8		READ2	REAL*8
	DUMP	REAL*8		SUHCAL	REAL*8
	etom1	REAL*8		TOFEP	REAL*8
	init2	REAL#8		WBBAL	REAL*8
	READ1	REAL*8		WELLSS	REAL*8
	READ2	REAL#8		WELRIS	REAL*8
	SUMCAL	REAL#8		WRITE2	REAL*8
	WELLSS	REAL*8		WRITE5	REAL*8
	WELRIS	REAL#8			
	WRITE2	REAL*8	PAEHDT	HST3D	REAL=8 DIMENSION(10
				BLOCKDATA	REAL*8 DIMENSION(10
P1	PRNTAR	CHARACTER*2		DUMP	REAL*8 DIMENSION(10
				INIT2	REAL*8 DIMENSION(10
P2	PRNTAR	CHARACTER*2		READ1	REAL*8 DIMENSION(10
				TOFEP	REAL*8 DIMENSION(10
P4	PRNTAR	CHARACTER*2			
			PAIF	APLYBC	REAL*8 DIMENSION(*)
P5	PRNTAR	CHARACTER*2		INIT2	REAL*8 DIMENSION(*)

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Table	11.2	2Cross-reference	list of	f variab.	lesContinued
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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
PARA	REWI3	REAL*8 DIMENSION(*)	PLBL	WRITE5	CHARACTER*20
PARB	REWI3	REAL*8 DIMENSION(*)	PLOTWC	HST3D	LOGICAL*4
PARC	REWI3	REAL*8 DIMENSION(*)		ERROR'S PLOTOC	LOGICAL*4 LOGICAL*4
PCL	WELLSS	REAL*8		READ3	LOGICAL*4
		· · · · ·	PLOTWP	HST3D	LOGICAL#4
PCR	WELLSS	REAL*8		ERROR3	LOGICAL#4
				PLOTOC	LOGICAL*4
PCS	PLOTOC	REAL*8 DIMENSION(*)		READ3	LOGICAL*4
PCW	PLOTOC	REAL*8 DIMENSION(*)	PLOTWT	HST3D	LOGICAL*4
				ERROR3	LOGICAL*4
PFAC	MAP2D	REAL*8		PLOTOC	LOGICAL*4
PFSLOW	SUMCAL	LOGICAL*4		READ3	LOGICAL*4
			PLTZON	HST3D	LOGICAL#4
PHILBC	APLYBC	REAL*8 DIMENSION(*)		DUMP	LOGICAL*4
	ERROR3	REAL*8 DIMENSION(*)		READI	LOGICAL*4
	INIT3	REAL*8 DIMENSION(*)		READ2	LOGICAL*4
	WRITE3	REAL*8 DIMENSION(*)		WRITE2	LOGICAL#4
	•	. • •		WRITE3	LOGICAL*4
PI	APLYBC	REAL*8			<i>x</i>
· •	INIT2	REAL*8	PLUS	PLOT	CHARACTER*1
	WELLSS	REAL*8			
	WELRIS	REAL*8	PMCHDT	CALCC	REAL*8
	WFDYDZ	REAL*8			· · · ·
		5547.40	PMCHV	CALCC	REAL*8
PINIT	HST3D	REAL*8		INIT2	REAL*8 DIMENSION(*
	DUMP	REAL*8		SUMCAL	REAL*8 DIMENSION(*
	ETOM1	REAL*8		•	
	INIT2	REAL*8	PMCV	CALCC	REAL*8
	READ1	REAL*8		init2	REAL*8 DIMENSION(*
	READ2	REAL*8		SUMCAL	REAL*8 DIMENSION(*
	WRITE2	REAL*8			

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
PMCVDT	CALCC	REAL*8	PPCR	WELLSS	REAL*8
PMHV	CALCC	REAL*8	PRBCF	HET3D	LOGICAL*4
	INIT2	REAL*8 DIMENSION(*)		SUMCAL	LOGICAL*4
	SUHCAL	REAL*8 DIMENSION(*)		WRITE5	LOGICAL*4
PMHVDT	CALCC	REAL*8	PRDV	HST3D	LOGICAL*4
	-			SUMCAL	LOGICAL*4
PNP	CALCC	REAL#8		WRITE5	LOGICAL*4
	ETOM2	REAL*8 DIMENSION (*)			
	INIT3	REAL*8 DIMENSION(*)	PRESS	TOPEP	REAL*8
	READ3	REAL*8 DIMENSION(*)		•	
			PRGFB	HST3D	LOGICAL*4
POROAR	HST3D	REAL*8		SUMCAL	LOGICAL*4
	APLYBC	REAL#8		WRITE5	LOGICAL*4
	DUMP	REAL*8			
	ETOM1	REAL*8	PRIBCY	HST3D	INTEGER*4
	INIT2	REAL*8		READ3	INTEGER*4
	READI	REAL*8		WRITES	INTEGER*4
	READ2	REAL*8			
	WRITE2	REAL*8	PRIDV	HST3D	INTEGER*4
			•	READ3	INTEGER*4
POROS	COEFF	REAL*8 DIMENSION(*)		WRITES	INTEGER*4
•••••	INIT2	REAL*8 DIMENSION(*)		<b>-</b> .	
	READ2	REAL *8 DIMENSION (*)	PRICFB	HST3D	INTEGER*4
	WRITE2	REAL*8 DIMENSION(*)		READ3	INTEGER*4
				WRITES	INTEGER*4
POS	PLOTOC	REAL*8 DIMENSION(*)			
			PRIKD	HST3D	INTEGER*4
Posup	HAP2D	LOGICAL*4		READ3	INTEGER#4
				WRITE4	INTEGER*4
Pow	PLOTOC	REAL*8 DIMENSION(*)		WRITE5	INTEGER*4
PPCL	WELLSS	REAL*8	PRIMAP	HST3D	INTEGER*4
				READ3	INTEGER*4
,				WRITE3	INTEGER*4
			1	<b>WRITES</b>	INTEGER*4
			/		

Table 11.2--Cross-reference list of variables--Continued

	Refer-			Refer-	÷ .
/ariable	encing	Variable	Variable	encing	Variable
name	programs	type	name	programs	type
PRIPTC	HST3D	INTEGER*4	PRTBC	HST3D	LOGICAL*4
	READ3	INTEGER#4		DUMP	LOGICAL*4
	WRITE5	INTEGER*4		READI	LOGICAL#4
				READ2	LOGICAL#4
PRISLM	HST3D	INTEGER*4		READ3	LOGICAL*4
	READ3	INTEGER*4		SOR2L	LOGICAL*4
	WRITE5	INTEGER*4		WELLSS	LOGICAL#4
				WELRIS	LOGICAL*4
Privel	HST3D	INTEGER*4		WRITE2	LOGICAL#4
	READ3	INTEGER#4		WRITE3	LOGICAL*4
9	WRITE4	INTEGER#4	,		
	WRITE5	INTEGER#4	PRTCCM	HST3D	LOGICAL*4
				DUMP	LOGICAL#4
Priwel	HST3D	INTEGER#4		READ1	LOGICAL#4
	READ3	INTEGER*4		READ2	LOGICAL#4
	WRITE5	INTEGER*4		SUMCAL	LOGICAL#4
RKD	WRITE4	LOGICAL#4	PRTCHR	MAP2D	CHARACTER*125
RNT	WRITE3	LOGICAL#4	PRTDV	HST3D	LOGICAL#4
				DUMP	LOGICAL#4
PRPTC	HST3D	LOGICAL*4		READ1	LOGICAL*4
	SUMCAL	LOGICAL*4		READ2	LOGICAL*4
	WRITE5	LOGICAL*4		WRITE2	LOGICAL#4
		•, •		WRITE3	LOGICAL#4
RSLM -	HST3D	LOGICAL*4		-	
	SUMCAL	LOGICAL*4			
1.4	WRITE5	LOGICAL#4			

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
PRTFP	HST3D	LOGICAL#4	PRTSLM	HST3D	LOGICAL=4
	DUHP	LOGICAL#4		DUHP	LOGICAL*4
	READ	LOGICAL#4		READ1	LOGICAL#4
	READ2	LOGICAL#4		READ2	LOGICAL#4
	WRITE2	LOGICAL#4		READ3	LOGICAL#4
	WRITE3	LOGICAL#4		SOR2L	LOGICAL*4
				WELLSS	LOGICAL#4
PRTIC	HST3D	LOGICAL#4		WELRIS	LOGICAL#4
	DUMP	LOGICAL=4		WRITE2	LOGICAL*4
	READI	LOGICAL#4		WRITE3	LOGICAL*4
	READ2	LOGICAL#4			
	WRITE2	LOGICAL#4	PRTWEL	hs <b>t3</b> d	LOGICAL*4
	WRITE3	LOGICAL=4		DUMP	LOGICAL*4
				READ1	LOGICAL*4
PRTHPD	HST3D	LOGICAL=4		READ2	LOGICAL*4
	READ3	LOGICAL#4		READ3	LOGICAL#4
	WRITE3	LOGICAL#4		WELLSS	LOGICAL*4
	WRITE5	LOGICAL#4		WELRIS	LOGICAL*4
				WRITE2	LOGICAL#4
PRTPHP	HST3D	LOGICAL#4		WRITE3	LOCICAL*4
	DUMP	LOGICAL#4		_	
	READI	LOGICAL#4	PRVEL	WRITE4	LOGICAL*4
	READ2	LOGICAL#4			
	WRITE2	LOGICAL#4	PRWEL	HST3D	LOGICAL#4
	WRITE3	LOGICAL#4		SUHCAL	LOGICAL*4
				WRITES	LOGICAL*4
PRTRE	hst3d	LOGICAL#4			
	DUMP	LOGICAL#4	PSBC	ASEMBL	REAL*8 DIMENSION(*)
	IREWI	LOGICAL*4		INIT3	REAL*8 DIMENSION(*)
	READ1	LOGICAL*4		WRITE3	REAL*8 DIMENSION(*)
	READ2	LOGICAL*4			
	REWI	LOGICAL#4	PSLBL	PLOTOC	CHARACTER*50
	REWI 3	LOGICAL*4			
	SUMCAL	LOGICAL*4	C.		

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
PSMAX	PLOTOC	REAL*8	PWLBL	PLOTOC	CHARACTER*50
PSMIN	PLOTOC	REAL*8	PWMAX	PLOTOC	REAL*8
PTOP	WELLSS	REAL*8	PWMIN	PLOTOC	REAL*8
PU	APLYBC	REAL*8	PWREND	HST3D	REAL*8
				WBBAL	REAL*8
PV	CALCC	REAL*8		WELLSS	REAL*8
	COEFF	REAL*8 DIMENSION(*)		WELRIS	REAL*8
	INIT2	REAL*8 DIMENSION(*)			
	SUMCAL	REAL*8 DIMENSION(*)	PWRK	WFDYDZ	REAL*8
. ·	WRITE2	REAL*8 DIMENSION(*)			
			PWSUR	WBBAL	REAL*8 DIMENSION(*)
PVDTN	CALCC	REAL*8		WELLSS	REAL*8 DIMENSION(*)
				WRITE5	REAL*8 DIMENSION(*)
PVK	CALCC	REAL*8			
i	COEFF	REAL*8 DIMENSION(*)	PWSURS	ERROR3	REAL*8 DIMENSION(*)
	INIT2	REAL*8 DIMENSION(*)	,	READ3	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8 DIMENSION(*)		WELLSS	REAL*8 DIMENSION(*)
		•		WRITE3	REAL*8 DIMENSION(*)
PVKDTN	CALCC	REAL*8	•		
	•	•	QDVSBC	SBCFLO	REAL*8 DIMENSION(*)
PWCELL	WRITE5	REAL*E			
			QFAC	MAP2D	REAL <sup>48</sup>
PWKT	ASEMBL	REAL*8 DIMENSION(*)	-		
	ERROR3	REAL*8 DIMENSION(*)	QFAIF	SUMCAL	REAL*8 DIMENSION(*)
	ITER	REAL*8 DIMENSION(*)	•	WRITE5	REAL*8 DIMENSION(*)
	READ3	REAL*8 DIMENSION(*)			
	WBBAL	REAL*8 DIMENSION(*)	QFBC	APLYBC	REAL <sup>+8</sup>
	WELLSS	REAL*8 DIMENSION(*)	•	ASEMBL	REAL*8
	WRITE3	REAL*8 DIMENSION(*)		SUMCAL	REAL*8
	WRITE5	REAL*8 DIMENSION(*)		•	
PWKTS	ITER	REAL*8 DIMENSION(*)			
	WBBAL	REAL*8 DIMENSION(*)			

REAL\*8 DIMENSION(\*)

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
QFBCV	APLYBC	REAL*8 DIMENSION(*)	QHCBC	APLYBC	REAL*8 DIMENSION(*)
•	ASEMBL	REAL+8 DIMENSION(*)	-	SUMCAL	REAL*8 DIMENSION(*)
	ERROR3	REAL*8 DIMENSION(*)		WRITE5	REAL*8 DIMENSION(*)
	INIT3	REAL*8 DIMENSION(*)			
	ITER	REAL*8 DIMENSION (*)	QHFAC .	HST3D	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)	-	WELRIS	REAL#8
	WRITE3	REAL*8 DIMENSION(*)		WFDYDZ	REAL*8
	WRITES	REAL*8 DIMENSION (*)			
		•••	<b>QHFBC</b>	APLYBC	REAL*8 DIMENSION(*)
QFFX	ETOM2	REAL*8 DIMENSION(*)	•	ERROR3	REAL*8 DIMENSION(*)
•	INIT3	REAL*8 DIMENSION(*)		INIT3	REAL*8 DIMENSION(*)
	READ3	REAL*8 DIMENSION(*)		SUNCAL	REAL*8 DIMENSION(*)
				WRITE3	REAL*8 DIMENSION(*)
qffy	ETOM2	REAL*8 DIMENSION(*)		WRITE5	REAL*8 DIMENSION(*)
	init3	REAL*8 DIMENSION(*)			
	READ3	REAL*8 DIMENSION(*)	QHFX	ETON2	REAL*8 DIMENSION(*)
	•		•	INIT3	REAL*8 DIMENSION(*)
QFFZ	etom2	REAL*8 DIMENSION(*)		READ3	REAL*8 DIMENSION (*)
•	init3	REAL*8 DIMENSION(*)			
	READ3	REAL*8 DIMENSION(*)	QHFY	ETOM2	REAL*8 DIMENSION(*)
			•	INIT3	REAL*8 DIMENSION(*)
QFLBC	SUMCAL	REAL*8 DIMENSION(*)		READ3	REAL*8 DIMENSION(*)
•	WRITE5	REAL*8 DIMENSION(*)			x
			QHFZ	etom2	REAL*8 DIMENSION(*)
QFSBC	ASEMBL	REAL*8 DIMENSION(*)	-	INIT3	REAL*8 DIMENSION(*)
	ITER	REAL*8 DIMENSION(*)		READ3	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8 DIMENSION(*)			
	WRITES	REAL*8 DIMENSION(*)	QHLBC	SUMCAL	REAL*8 DIMENSION(*)
			_ *	WRITE5	REAL*8 DIMENSION(*)
QHAIF	SUMCAL	RÉAL*8 DIMENSION(*)			
-	WRITE5	REAL*8 DIMENSION(*)	QHLYR	WBBAL	REAL*8 DIMENSION(*)
				WELLSS	REAL*8 DIMENSION(*)
QHBC	APLYBC	REAL*8		WRITE5	REAL*8 DIMENSION(*)
-	ASEMBL	REAL*8			

Table 11.2--Cross-reference list of variables--Continued

	Refer-	• :		Refer-	<b></b>
Variable	encing	Variable	Variable	encing	Variable
name	programs	ams type	name	programs	type
QHSBC	SUMCAL	REAL*8 DIMENSION(*)	QSFX	ETOM2	REAL*8 DIMENSION(*)
•	WRITE5	REAL*8 DIMENSION(*)	·	INIT3	REAL*8 DIMENSION(*)
		•		READ3	REAL*8 DIMENSION(*)
QHW	ASEMBL	REAL*8			
•	SUMCAL	REAL*8 DIMENSION(*)	QSFY	ETOM2	REAL*8 DIMENSION(*)
	WBBAL	REAL*8 DIMENSION(*)		INIT3	REAL*8 DIMENSION(*)
	WELLSS	REAL*8 DIMENSION(*)		READ3	REAL*8 DIMENSION(*)
	WRITES	REAL*8 DIMENSION(*)			
	2		QSF2	ETOM2	REAL*8 DIMENSION(*)
QHWRK	WFDYDZ	REAL*8		INIT3	REAL*8 DIMENSION(*)
•				READ3	REAL*8 DIMENSION(*)
QLIM	APLYBC	REAL*8			
•	ASEMBL	REAL*8	QSLBC	SUMCAL	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8		WRITE5	REAL*8 DIMENSION(*)
QN	APLYBC	REAL*8	QSLYR	WBBAL	REAL*8 DIMENSION(*)
•••	ASEMBL	REAL*8		WELLSS	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8	- u	WRITE5	REAL*8 DIMENSION(*)
QNP	ASEMBL	REAL*8	QSSBC	SUMCAL	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8		WRITE5	REAL*8 DIMENSION(*)
QSAIF	SUMCAL	REAL*8 DIMENSION(*)	qsw	ASEMBL	REAL*8
<b>.</b>	WRITE5	REAL*8 DIMENSION(*)		SUMCAL	REAL*8 DIMENSION(*)
				WBBAL	REAL*8 DIMENSION(*)
QSBC	APLYBC	REAL*6	· · ·	WELLSS	REAL*8 DIMENSION(*)
•	ASEMBL	REAL*8		WRITE5	REAL*8 DIMENSION(*)
QSFBC	APLYBC	REAL*8 DIMENSION(*)	QUOT	BSODE	REAL*8 DIMENSION(11,2)
<b>.</b>	ERROR3	REAL*8 DIMENSION(*)	•	•	
	INIT3	REAL*8 DIMENSION(*)	QUOTSV	BSODE	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)			•
	WRITE3	REAL*8 DIMENSION(*)	QWAV	ASEMBL	REAL*8
	WRITE5	REAL*8 DIMENSION(*)	-		

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
QWLYR	ASEMBL	<b>REAL*8</b> DIMENSION(*)	RBW	D4DES	INTEGER*4 DIMENSION(*
	ITER	REAL*8 DIMENSION(*)		ORDER	INTEGER*4 DIMENSION(*
	WBBAL	REAL*8 DIMENSION(*)			
	WBCFLO	REAL*8 DIMENSION(*)	RCPPM	etom1	REAL*8 DIMENSION(*)
	WELLSS	REAL*8 DIMENSION(*)		INIT2	REAL*8 DIMENSION(*)
	WRITES	REAL*8 DIMENSION(*)		READ2	REAL*8 DIMENSION(*)
				WRITE2	REAL*8 DIMENSION(*)
QWM	ASEMBL	REAL*8 DIMENSION(*)			
-	ITER	REAL*8 DIMENSION(*)	RDAIF	hst3d	LOGICAL*4
	SUMCAL	REAL*8 DIMENSION(*)		ERROR3	LOGICAL*4
	WBBAL	REAL*8 DIMENSION(*)		etom2	LOGICAL*4
	WELLSS	REAL*8 DIMENSION(*)		INI <b>T</b> 3	LOGICAL*4
	WELRIS	REAL#8		READ3	LOGICAL*4
	WRITES	REAL*8 DIMENSION(*)		WRITE3	LOGICAL*4
QWN	ASEMBL	REAL*8	RDCALC	HST3D	LOGICAL=4
-				ERROR3	LOGICAL#4
QWNP	ASEMBL	REAL*8		ETOM2	LOGICAL*4
•				INIT3	LOGICAL*4
QWR	HST3D	REAL*8		READ3	LOGICAL+4
•	WELRIS	REAL#8		WRITE3	LOGICAL*4
	WFDYDZ	REAL*8			
	•		RDECHO	HST3D	LOGICAL*4
QWV	ASEMBL	REAL*8 DIMENSION(*)		CLOSE	LOGICAL*4
•	ERROR3	REAL*8 DIMENSION(*)		DUMP	LOGICAL*4
	READ3	REAL*8 DIMENSION(*)		IREVI	LOGICAL*4
	SUNCAL	REAL*8 DIMENSION(*)		PLOTOC	LOGICAL*4
	WBBAL	REAL*8 DIMENSION(*)		READ1	LOGICAL*4
	WELLSS	REAL*8 DIMENSION(*)		READ2	LOGICAL*4
	WRITE3	REAL*8 DIMENSION(*)		READ3	LOGICAL*4
	WRITES	REAL*8 DIMENSION(*)		REWI	LOGICAL*4
R	WELLSS	REAL*8 DIMENSION(*)			

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Variable na <del>me</del>	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
RDFLXH	HST3D	LOGICAL*4	RDPLTP	PLOTOC	LOGICAL*4
	ERROR3	LOGICAL*4			
	etom2	LOGICAL#4	RDSCBC	HST3D	LOGICAL*4
	INIT3	LOGICAL#4		ERROR3	LOGICAL*4
	READ3	LOGICAL#4		ETOM2	LOGICAL*4
	WRITE3	logical#4		INIT3	LOGICAL*4
		• •		READ3	LOGICAL*4
RDFLXQ .	RST3D	LOGICAL#4		WRITE3	LOGICAL*4
	ERROR3	LOGICAL#4			
	ETOM2	LOGICAL#4	RDSPBC	HST3D	LOGICAL*4
	INIT3	LOGICAL#4		ERROR3	LOGICAL#4
	READ3	LOGICAL*4		ETOM2	LOGICAL*4
	WRITE3	LOGICAL*4		INIT3	LOGICAL*4
				READ3	LOGICAL*4
RDFLXS	HST3D	LOGICAL#4		WRITE3	LOGICAL*4
	ERROR3	LOGICAL*4		_	
	ETOM2	LOGICAL#4	EDSTEC	HST3D	LOGICAL*4
	INIT3	LOGICAL*4		ERROR3	LOGICAL#4
	READ3	LOGICAL#4		ETOM2	LOGICAL#4
	WRITE3	LOGICAL*4		init3	LOGICAL*4
				READ3	LOGICAL*4
RDLBC	HST3D	LOGICAL#4		WRITE3	LOGICAL*4
	ERROR3	LOGICAL*4			
	ETOM2	LOGICAL*4	RDVAIF	HST3D	LOGICAL*4
	INIT3	LOGICAL*4		APLYBC	LOGICAL*4
	READ3	Logical*4		DUMP	LOGICAL*4
	WRITE3	LOGICAL*4		ERROR2	LOGICAL*4
				ETOM1	LOGICAL*4
RDMPDT	HST3D	LOGICAL*4		INIT2	LOGICAL#4
	ERROR3	LOGICAL*4		READ1	LOGICAL*4
	ETOM2	LOGICAL*4		READ2	LOGICAL*4
	INIT3	LOGICAL*4		WRITE2	LOGICAL*4
	READ3	LOGICAL*4			
	WRITE3	LOGICAL*4			

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
RDWDEF	HST3D	LOGICAL*4	RF	APLYBC	REAL*8 DIMENSION(*)
	DUMP	LOGICAL*4		ASEMBL	REAL*8 DIMENSION(*)
	ETOM1	LOGICAL*4		COEFF	REAL*8 DIMENSION(*)
	INIT2	LOGICAL*4		ITER	REAL*8 DIMENSION(*)
	ITER	LOGICAL*4		WELLSS	REAL*8 DIMENSION(*)
	READ1	LOGICAL*4			
	READ2	LOGICAL*4	RH	APLYBC	REAL*8 DIMENSION(*)
	WBBAL	LOGICAL#4		ASEMBL	REAL*8 DIMENSION(*)
	WELLSS	LOGICAL*4		COEFF	REAL*8 DIMENSION(*)
	WELRIS	LOGICAL*4		ITER	REAL*8 DIMENSION(*)
	WRITE2	LOGICAL*4		WELLSS	REAL*8 DIMENSION(*)
	WRITE5	LOGICAL*4			
			RH1	ASEMBL	REAL*8 DIMENSION(*)
RDWFLO	HST3D	LOGICAL#4			
	ERROR3	LOGICAL*4	RHS	ASEMBL	REAL*8 DIMENSION(*)
	ETOM2	LOGICAL#4		D4DES	REAL*8 DIMENSION(*)
	INIT3	LOGICAL*4		iter	REAL*8 DIMENSION(*)
	READ3	LOGICAL#4		L2SOR	REAL*8 DIMENSION(*)
	WRITE3	LOGICAL*4		SOR2L	REAL*8 DIMENSION(*)
RDWHD	HST3D	LOGICAL*4	RHSSBC	ASEMBL	REAL*8 DIMENSION(*)
	ERROR3	LOGICAL#4		SECFLO	REAL*8 DIMENSION(*)
	ETOM2	LOGICAL#4			
	INIT3	LOGICAL*4	RHSW	ASEMBL	REAL*8 DIMENSION(*)
	READ3	LOGICAL#4		WBCFLO	REAL*8 DIMENSION(*)
	WRITE3	LOGICAL#4			
			RIOAR	HST3D	REAL*8
REN	WELLSS	REAL*8		APLYBC	REAL*8
	WFDYDZ	REAL*8		DUMP	REAL*8
				ETOM1	REAL*8
RESTRT	HST3D	LOGICAL*4		INIT2	REAL*8
	READI	LOGICAL*4		READI	REAL*8
	WRITE1	LOGICAL*4		/ READ2	REAL*8
				WRITE2	REAL*8

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
RM	INIT2	REAL*8 DIMENSION(*)	SDECAY	HST3D	REAL*8
	WRITE2	REAL*8 DIMENSION(*)		SUMCAL WRITES	REAL*8 REAL*8
RORW2	INIT2	REAL*8		WKLIEJ	KEAL-O
			SHRES	HST3D	REAL*8
RPRN	PRNTAR	CHARACTER*1	•	SUMCAL	REAL*8
				WRITE5	REAL*8
RS	APLYBC	REAL*8 DIMENSION(*)			
	ASEMBL	REAL*8 DIMENSION(*)	SHRESF	HST3D	REAL*8
	COEFF	REAL*8 DIMENSION(*)		SUMCAL	REAL*8
	ITER	REAL*8 DIMENSION(*)		WRITE5	REAL*8
	WELLSS	REAL*8 DIMENSION(*)			
			SIR	HST3D	REAL*8
RSI ASI	ASEMBL	REAL*8 DIMENSION(*)		SUMCAL	REAL*8
				WRITE5	REAL*8
SAVLDO	HST3D	Logical#4			1
	CLOSE	LOGICAL*4	SIRO	HST3D	REAL*8
	DUMP	LOGICAL*4		APLYBC	REAL*8
	READ3	LOGICAL*4		DUMP	REAL*8
				INIT2	REAL*8
SCALMF	HST3D	LOGICAL*4		READ1	REAL*8
	CLOSE	LOGICAL*4		SUMCAL	REAL*8
	DUMP	LOGICAL*4		WRITE2	REAL#8
	INIT2	LOGICAL*4		WRITE5	REAL®8
	IN1T3	LOGICAL#4			
	IREWI	LOGICAL*4	SIRN	SUMCAL	REAL*8
	PLOTOC	LOGICAL#4			
	READ1	LOGICAL*4			
	READ2	LOGICAL#4		-	
	READ3	LOGICAL*4	i.		
	REWI	LOGICAL#4			
	SUNCAL	LOGICAL <sup>4</sup> 4		•	
	WRITE1	LOGICAL#4			
	WRITE2	LOGICAL*4			
	WRITE3	LOGICAL*4			

Table 11.2--Cross-reference list of variables--Continued

WRITE5

LOGICAL\*4

Variable	Refer- encing	Variable	Variable	Refer- eucing	Variable
name	programs	type	name	programs	type
SLMETH	HST3D	INTEGER#4	SOLUTE	HST3D	LOGICAL*4
	ASEMBL	Integer+4		APLYBC	LOGICAL*4
	DUHP	INTEGER#4		ASEMBL	LOGICAL*4
	ERROR1	INTEGER#4		CALCC	LOGICAL*4
	init1	INTEGER#4		COEFF	LOGICAL*4
	INIT2	Integer+4		CRSDSP	LOGICAL*4
	ITER	Intecer*4		DUMP	LOGICAL#4
	READ1	INTEGER#4		ERROR2	LOGICAL*4
SBCF1 WBCF1	READ2	INTEGER*4		ERROR3	LOGICAL*4
	SBCFLO	INTECER#4		etom1	LOGICAL#4
	WBCFLO	Intecer+4		INITL	LOGICAL*4
	WRITE1	Integer#4		INIT2	LOGICAL*4
	WRITE2	INTEGER*4		INIT3	LOGICAL*4
	WRITES	1nteger+4		ITER	LOGICAL*4
				PLOTOC	LOGICAL*4
SHCALC	CALCC	LOGICAL#4		READ1	LOGICAL*4
				READ2	LOGICAL*4
				READ3	LOGICAL*4
				SUMCAL	LOGICAL*4
				VISCOS	LOGICAL*4
				WBBAL	LOGICAL*4
				WELLSS	LOGICAL*4
				WRITE1	LOGICAL#4
				WRITE2	LOGICAL*4
				WRITE3	LOGICAL*4
				WRITE4	LOGICAL*4
				WRITE5	LOGICAL*4

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
SOLVE	L2SOR	LOGICAL*4	STOTFP	HST3D	REAL*8
	SOR2L	LOGICAL*4	•	APLYBC	REAL*8
				SUMCAL	REAL*8
SPR	HST3D	REAL*8 DIMENSION(3)		WRITE5	REAL*8
	L2SOR	REAL*8 DIMENSION(3)			· · · ·
•	00000		STOTHI	HST3D	REAL*8
SPRAD	L2SOR	REAL*8	0.000	APLYBC	REAL*8
	SOR2L	REAL*8		SUMCAL	REAL*8
	Dollar			WRITE5	REAL*8
SRES	HST3D	REAL*8			
·	SUMCAL	REAL*8	STOTHP	HST3D	REAL*8
· ·	WRITE5	REAL*8	0101	APLYBC	REAL*8
				SUMCAL	REAL*8
SRESF	HST3D	REAL*8		WRITE5	REAL*8
	SUMCAL	REAL*8			
	WRITES	REAL*8	STOTSI	HST3D	REAL*8
			0.000	APLYBC	REAL*8
SSRES	HST3D	REAL#8		SUMCAL	REAL*8
bbillip	SUMCAL	REAL*8		WRITE5	REAL*8
	WRITE5	REAL*8			
			STOTSP	HST3D	REAL*8
SSRESF	HST3D	REAL*8	010101	APLYBC	REAL*8
001001	SUMCAL	REAL*8		SUMCAL	REAL*8
	WRITE5	REAL*8		WRITE5	REAL*8
STOTFI	HST3D	REAL*8	SUM	INIT2	REAL*8
-	APLYBC	REAL*8			• • • •
	SUMCAL	REAL*8	SUM1	ITER	REAL*8
	WRITE5	REAL*8		WELLSS	REAL*8

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
SUM2	ITER	REAL*8	SZZM	ASEMBL	REAL*8
Sumcui	WELLSS	REAL*8	SZZP	ASEMBL	REAL=8
SUMMOB	WELLSS	REAL*8	SZZW	WELLSS	REAL*8
Sumtwi	WELLSS	REAL#8	T	APLYBC	REAL*8 DIMENSION(*)
				ASEMBL	REAL*8 DIMENSION(*)
Sumwi	WELLSS	REAL*8		CALCC	REAL*8
				COEFF	REAL*8 DIMENSION(*)
SUHCCE	VSINIT	REAL*8		CRSDSP	REAL*8 DIMENSION(*)
				ÉTOM1	REAL*8 DIMENSION(*)
SUMXY	VSINIT	.REAL*8		INIT2	REAL*8 DIMENSION(*)
				Iter	REAL*8 DIMENSION(*)
SVBC	ASEMBL	LOGICAL*4		READ2	REAL*8 DIMENSION(*)
				SUNCAL	REAL*8 DIMENSION(*)
SXX	ASEMBL	REAL*8 DIMENSION(*)		TOFEP	REAL*8
	COEFF	REAL*8 DIMENSION(*)		VISCOS	REAL*8
				WELLSS	REAL*8 DIMENSION(*)
MXX	ASEMBL	REAL*8		WRITE2	REAL*8 DIMENSION(*)
		· ·		WRITES	REAL*8 DIMENSION (*)
SXXP	ASEMBL	REAL*8			•••
		•	TO	HST3D	REAL*8
SYMID	MAP2D	CHARACTER*1 DIMENSION(5)	-	DUMP	REAL*8
				ETOH1	REAL*8
SYY	ASEMBL	REAL*8 DIMENSION(*)		INIT2	REAL*8
	COEFF	REAL*8 DIMENSION (*)		READI	REAL*8
		•••••		READ2	REAL*8
SYYM	ASEMBL	REAL*8		SUNCAL	REAL*8
				WBBAL	REAL*8
SYYP	ASEMBL	REAL*8		WELLSS	REAL*8
				WFDYDZ	REAL*8
SZZ	ASEMBL	REAL*8 DIMENSION(*)		WRITE2	REAL*8
, a,	COEFF	REAL*8 DIMENSION(*)		WRITE5	REAL*8

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable Lype
T00	HST3D	REAL*8	TAMBK	HSTJD	REAL*8
	WBBAL	REAL*8		WELRIS	REAL*8
	WELLSS	REAL*8		WFDYDZ	REAL*8
· .	WELRIS	REAL*8			
			TATWR	ETOM1	REAL*8 DIMENSION(*)
TOH	HST3D	REAL*8		READ2	REAL*8 DIMENSION(*)
-	APLYBC	REAL*8		WELRIS	REAL*8
	ASEMBL	REAL+8		WRITE2	REAL*8 DIMENSION(*)
	DUMP	REAL*8			
	ETOH1	REAL*8	TC	PLOTOC	REAL*8 DIMENSION(*)
	INIT2	REAL*8			• •
	READ1	REAL#8	TCS	PLOTOC	REAL*8 DIMENSION(*)
	READ2	REAL*8		•	
	SUMCAL	REAL*8	TCW	PLOTOC	REAL*8 DIMENSION(*)
2	WELLSS	REAL*8			
	WELRIS	REAL*8	TDATA	VSINIT	REAL*8 DIMENSION(*)
	WRITE2	REAL*8			
	WRITE5	REAL*8	TDEHIR	WRITE5	REAL*8
TA	BSODE	REAL*8	TDFIR	WRITE5	REAL*8
TABWR	etom1	REAL*8 DIMENSION(*)	TDSIR	WRITE5	REAL#8
	READ2	REAL*8 DIMENSION(*) REAL*8	TDX	COEFF	REAL*8
	WELRIS	REAL*8 DIMENSION(*)	102	WVLF F	pitana U
	WRITE2	VEUT-O NTURUSIAN(.)	TDXY	COEFF	REAL*8
TAIF	APLYBC	REAL*8 DIMENSION(*)	Thur	VULLE	ay setting " " W
TUTE	ASEMBL	REAL*8 DIMENSION(*)	TDXZ	COEFF	REAL*8
	INIT3	REAL*8 DIMENSION(*)	LUAG	uvge e	ALLER BY THE
	SUMCAL	REAL*8 DIMENSION(*)	TDY	COEFF	REAL*8
	JUNUAL	MERIN D REIMARKAUL	141	ANNE L	evneting - W
TAMBI	HST3D	REAL*8	TDYX	COEFF	REAL*8
	WELRIS	REAL*8	4 44 6 FS	www.h	
	WFDYDZ	REAL*8	TDYZ	COEFF	REAL*8
	*** L/ L L/ 60				

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs		riable type	Variable name	Refer- encing programs	v	ariable type
TDZ	COEFF	REAL*8		TFRESP	HST3D	REAL*8	
					SUMCAL	REAL*8	
TDZX	COEFF	REAL*8			WRITE5	REAL*8	
TDZY	COEFF	REAL*8		TFW	ASEMBL		DIMENSION(*)
					WELLSS	REAL*8	DIMENSION(*)
TERDT	HST3D	REAL*8	DIMENSION(14)				
	BLOCKDATA		DIMENSION(14)	TFX	COEFF	REAL*8	DIMENSION(*)
	DUMP	REAL*8	DIMENSION(14)		WRITE4	REAL*8	DIMENSION(*)
	INIT2	REAL*8	DIMENSION(14)				
	READI	REAL*8	DIMENSION(14)	TFXM	ASEHBL	REAL*8	
	TOFEP	REAL*8	DIMENSION(14)				
				TFXP	ASEMBL	REAL*8	
Tehst	HST3D	REAL*8	DIMENSION(32)				
	BLOCKDATA	REAL#8	DIMENSION (32)	TFY	COEFF	REAL*8	DIMENSION(*)
	DUMP	REAL#8	DIMENSION(32)		WRITE4	REAL*8	DIMENSION(*)
	INIT2	REAL*8	DIMENSION(32)				
	READ1	REAL*8	DIMENSION(32)	TFYM	ASEMBL	REAL*8	
	TOFEP	REAL*8	DIMENSION(32)				
				TFYP	ASEMBL	REAL#8	
TEMP	TOPEP	REAL*8					
	•			TFZ	COEFF	REAL*8	DIMENSION(*)
TFLX	APLYBC	REAL*8	DIMENSION(*)		WRITE4		DIMENSION(*)
	INIT3	REAL*8	DIMENSION(*)				5.0
	SUMCAL		dimension(*)	TF2M	ASEMBL	REAL*8	
	WRITE3	REAL*8	DIMENSION(*)				
	WRITE5	REAL*8	DIMENSION(*)	TFZP	ASEMBL	REAL*8	
TFRES	HST3D	REAL*8		THCBC	APLYBC	REAL*8	DIMENSION(*)
	SUHCAL	REAL*8			INIT2	REAL*8	DIMENSION(*)
	WRITES	REAL*8			-		• -

Table 11.2--Cross-reference list of variables--Continued

Variable	Refer- encing	Variable	Variable	Refer- encing	Variable
name	programs	type	name	programs	type
THETXZ	HST3D	REAL*8	THRU	HST3D	LOGICAL*4
	DUMP	REAL*8		COEFF	LOGICAL*4
	ERROR2	REAL*8		READ3	LOGICAL*4
	INIT2	REAL*8		WRITE4	LOGICAL*4
	READ1	REAL#8			
	READ2	REAL*8	THX	COEFF	REAL*8 DIMENSION(*)
	WRITE2	REAL#8		WRITE4	REAL*8 DIMENSION(*)
TRETYZ	HST3D	REAL*8	THXM	ASEMBL	REAL*8
	DUMP	REAL*8			
	ERROR2	REAL <sup>48</sup>	THXP	ASEMBL	REAL*8
	INIT2	REAL*8			
	<b>READI</b>	REAL*8	THXY	COEFF	REAL*8 DIMENSION(*)
	READ2	REAL*8		CRSDSP	REAL*8 DIMENSION(*)
	WRITE2	REAL*8		WRITE4	REAL*8 DIMENSION(*)
THETZZ	HST3D	REAL*8	THXZ	COEFF	REAL+8 DIMENSION(+)
	DUMP	REAL*8		CRSDSP	REAL*8 DIMENSION(*)
	ERROR2	REAL*8		WRITE4	REAL*8 DIMENSION(*)
	INIT2	REAL*8			
	READ1	REAL*8	THY	COEFF	REAL*8 DIMENSION(*)
•	READ2	REAL*8		WRITE4	REAL*8 DIMENSION(*)
	WRITE2	REAL*8			· ·
			THYM	ASEMBL.	REAL#8
TRI	TOFEP	REAL*8		· .	
			THYP	ASEMBL	REAL#8
THRES	HST3D	REAL*8			
	SUHCAL	REAL*8	THYX	COEFF	REAL*8 DIMENSION (*)
· . ·	WRITE5	REAL*8	x	CRSDSP	REAL*8 DIMENSION(*)
		·.	•	WRITE4	REAL*8 DIMENSION(*)
THRESF	hstjd	REAL*8			· · · · · · · · · · · · · · · · · · ·
	SUMCAL	REAL*8	THYZ	COEFF	REAL*8 DIMENSION(*)
	WRITE5	REAL*8		CRSDSP	REAL*8 DIMENSION (*)
				WRITE4	REAL*8 DIMENSION(*)

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Table 11,2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
	brograms				-77-
THZ	COEFF	REAL*8 DIMENSION(*)	TIMDAY	WELLSS	REAL*8
	WRITE4	REAL*8 DIMENSION(*)			
			TIME	HST3D	REAL*8
TH2H	ASEMBL	REAL*8		APLYBC	REAL*8
				CALCC	REAL*8
THZP	ASEMBL	REAL#8		CLOSE.	REAL*8
				COEFF	REAL*8
THZX	COEFF	REAL*8 DIMENSION(*)		DUMP	REAL*8
	CRSDSP	REAL*8 DIMENSION(*)		ERROR3	REAL*8
	WRITE4	REAL*8 DIMENSION(*)		etom2	REAL*8
				INIT2	REAL*8
THZY	COEFF	REAL*8 DIMENSION(*)		INIT3	REAL*8
	CRSDSP	REAL*8 DIMENSION(*)		ITER	REAL*8
	WRITE4	REAL*8 DIMENSION(*)		READ1	REAL*8
				READ3	REAL*8
TILT	HST3D	LOGICAL*4		SUMCAL	REAL*8
	DUMP	LOGICAL*4		WBBAL	REAL*8
	ERROR2	LOGICAL#4		WELLSS	REAL*8
	INIT2	LOGICAL*4		WELRIS	REAL*8
	READ1	LOGICAL*4		WRITE2	REAL*8
	READ2	LOGICAL*4		WRITE3	REAL*8
	WRITE2	LOGICAL*4		VRITE4	REAL*8
				WRITES	REAL*8
TINCHC	HST3D	BEAL*8			,
	COEFF	REAL*8	TIMED	APLYBC	REAL*8
	DUMP	REAL*8		WELRIS	REAL*8
	ERROR3	REAL*8			
	ETOM2	REAL*8	TIMEDN	APLYBC	REAL*8
	INIT3	REAL*8			
	READ3	REAL*8	TIMRST	HST3D	REAL*8
	SUHCAL	REAL*8		READ1	REAL*8
	WRITE1	REAL*8		WRITE1	REAL*8
	WRITE3	REAL*8		•	
	WRITE4	REAL*8			
	WRITE5	REAL*8	1		

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable nam <del>e</del>	Refer- encing programs	Variable type
TITLE	HST3D	CHARACTER*160	TOLDEN	HST3D	REAL*8
	DUMP	CHARACTER*160		DUMP	REAL*8
	MAP2D	CHARACTER*80		INIT2	REAL*8
	READ1	CHARACTER*160		ITER	REAL*8
	WRITE1	CHARACTER*160		READI	REAL*8
	WRITE5	CHARACTER*80		READ2	REAL*8
				WRITE2	REAL*8
TITLEO	HST3D	CHARACTER*160		•	
•	READ1	CHARACTER*160	TOLDNC	HST3D	REAL*8
	WRITEL	CHARACTER*160		DUMP '	REAL*8
				INIT2	REAL*8
TLBC	APLYBC	REAL*8 DIMENSION(*)		ITER	REAL*8
	ASEMBL	REAL*8 DIMENSION(*)		READ1	REAL*8
	INIT3	REAL*8 DIMENSION(*)		READ2	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)			
	WRITE3	REAL*8 DIMENSION(*)	TOLDNT	HST3D	REAL*8
				DUMP	REAL*8
TLBL	WRITE5	CHARACTER*20		INIT2	REAL*8
				ITER	REAL*8
TLO	TOFEP	REAL*8		READ1	REAL*8
				READ2	REAL*8
TM1	SOR2L	REAL*8 DIMENSION(*)			
			TOLDPW	HST3D	REAL*8
TM2	SOR2L	REAL*8 DIMENSION(*)		DUMP	REAL*8
· ·				etom1	REAL*8
THLBL	PLOTOC	CHARACTER*30		INIT2	REAL*8
		+ 1		READ1	REAL*8
TNP	CALCC	REAL*8	•	READ2	REAL*8
	READ3	REAL*8 DIMENSION(*)		WELLSS	REAL*8
	<u> </u>			WRITE2	REAL*8
то	PLOTOC .	REAL*8 DIMENSION(*)			

Table 11.2--Cross-reference list of variables--Continued

	Refer-			Refer-		
Variable	encing	Variable	Variable	encing	Variable	
name	programs	type	name	programs	type	
TOLPPW	HST3D	REAL*8	TOTHI	HST3D	REAL*8	
	DUMP	REAL*8		APLYBC	REAL*8	
	etom1	REAL*8		DUMP	REAL*8	
	INIT2	REAL*8		INIT2	REAL*8	
	READ1	REAL*8		READ1	REAL*8	
	READ2	REAL*8		SUHCAL	REAL*8	
	WELLSS	REAL*8		WRITE5	REAL*8	
	WRITE2	REAL*8				
			TOTHP	HST3D	REAL*8	
tolqw	HST3D	REAL#8		APLYBC	REAL*8	
	DUMP	REAL*8		DUHP	REAL*8	
	etom1	REAL*8		init2	REAL*8	
	init2	REAL*8		READ1	REAL*8	
	READ1	REAL*8		SUMCAL	REAL*8	
	READ2	REAL*8		WRITE5	REAL*8	
	WELLSS	REAL#8				
	WRITE2	REAL*8	TOTSI	HST3D	REAL*8	
				APLYBC	REAL*8	
tos	PLOTOC	REAL*8 DIMENSION(*)		DUMP	REAL*8	
				INIT2	REAL*8	
totf1	hst3d	REAL*8		READ1	REAL*8	
	APLYBC	REAL*8		SUMCAL	REAL*8	
	DUMP	REAL*8		WRITE5	REAL*8	
	INIT2	REAL*8				
	READ 1	REAL*8	TOTSP	hst3d	REAL*8	
	SUHCAL.	REAL*8		APLYBC	REAL*8	
	WRITE5	REAL=8		DUMP	REAL*8	
				INIT2	REAL*8	
Totpp	hst3d	REAL*8		READ1	REAL*8	
	APLYBC	REAL*8		SUHCAL	REAL*8	
	DUHP	REAL*8		WRITE5	REAL*8	
	INIT2	REAL*8				
	READ1	REAL*8		•		
	SUHCAL	REAL*8				
	WRITE5	REAL*8				

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
TOTWFI	HST3D	REAL*8	TOTWSI	HST3D	REAL*8
	APLYBC	REAL*8		APLYBC	REAL*8
	DUMP	REAL*8		DUMP	REAL*8
•	INIT2	REAL#8		INIT2	REAL*8
	READI	REAL#8		READ1	REAL*8
	SUMCAL	REAL*8		SUMCAL	REAL*8
	WRITE5	REAL*8		WRITE5	REAL*8
TOTWFP	HST3D	REAL*8	TOTWSP	HST3D	REAL*8
•	APLYBC	REAL <sup>*8</sup>		APLYBC	REAL*8
	DUMP	REAL*8		DUMP	REAL*8
	INIT2	REAL*8		INIT2	REAL#8
	READ1	REAL*8		READ1	REAL#8
	SUHCAL	real*8		SUMCAL	REAL*8
	WRITES	REAL#8		WRITE5	REAL*8
TOTWHI	HST3D	REAL*8	TOW	PLOTOC	REAL*8 DIMENSION(*)
	APLYBC	REAL*8			
	DUMP	REAL*8	TP1	SOR2L	REAL*8 DIMENSION(*)
	INIT2	REAL#8			
	<b>READ1</b>	REAL*8	TP2	SOR2L	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8			
	WRITE5	REAL*8	TPHCBC	APLYBC	REAL*8 DIMENSION(*)
TOTWHP	HST3D	REAL*8	TQFAIF	HST3D	REAL*8
	APLYBC	REAL*8	-	APLYBC	REAL*8
	DUMP	REAL*8		DVMP	REAL*8
	INIT2	REAL*8		INIT2	REAL*8
	READ1	REAL*8		READI	REAL <sup>4</sup> 8
	SUMCAL	REAL*8		SUMCAL	REAL*8
	WRITE5	REAL*8		WRITE5	REAL*8

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Variable name	Refer- encing progr <del>ams</del>	Variable Lype	Variable name	Refer- encing programs	Variable type
TQFFBC	HST3D	REAL#8	TQHAIF	HST3D	REAL*8
•	APLYBC	REAL*8		APLYBC	REAL*8
	DUMP	REAL#8		DUMP	REAL*8
	INIT2	REAL*8		INIT2	REAL*8
	READ1	REAL*8		READ1	REAL*8
	SUNCAL	REAL+8		SUMCAL	REAL*8
	WRITE5	REAL#8		WRITES	REAL*8
TQFINJ	HST3D	REAL*8	TQHFBC	hst3d	REAL*8
-	SUMCAL	RKAL=8		APLYBC	REAL*8
	WELLSS	REAL#8		DUMP	REAL*8
	WRITE5	REAL#8		init2	REAL*8
				READ1	REAL*8
rqflbc 1	HST3D	REAL#8		SUHCAL	REAL*8
•	APLYBC	REAL#8		WRITES	REAL*8
	DUMP	REAL#8			
	init2	REAL#8	тоннвс	HST3D	REAL*8
	READ1	REAL#8		APLYBC	REAL*8
	SUMCAL	REAL#8		DUMP	REAL*8
	WRITE5	REAL#8		INIT2	REAL*8
				READ1	REAL*8
TQFPRO	HST3D	REAL#8		SUMCAL	REAL*8
	SUMCAL	REAL#8		WRITES	REAL*8
	WELLSS	REAL*8			
	WRITES	REAL*8	TQHINJ	hst3d	REAL*8
		· · · ·		SUHCAL	REAL*8
TQFSBC	HST3D	REAL*8		WELLSS	REAL*8
	APLYBC	REAL*8		WRITE5	REAL*8
	DUMP	REAL*8			
	INIT2	REAL*8			
	READ1	REAL*8			
	SUMCAL	REAL*8			
	WRITE5	REAL*8			

Table 11.2--Cross-reference list of variables--Continued

Variable	Refer- encing	Variable	Variable	Refer- encing	Variable
name	programs	type	name	programs	type
TQHLBC	HST3D	REAL*8	TQSINJ	HST3D	REAL*8
	APLYBC	REAL#8		SUMCAL	REAL*8
	DUMP	REAL*8		WELLSS	REAL*8
	INIT2	REAL#8	•	WRITE5	REAL*8
	READ1	REAL*8			· · · · · ·
	SUMCAL	REAL#8	TQSLBC	HST3D	REAL*8
	WRITE5	REAL*8		APLYBC	REAL*8
		-		DUMP	REAL*8
TQHPRO	HST3D	REAL*8		INIT2	REAL*8
-	SUMCAL	REAL*8		READ1	REAL*8
	WELLSS	REAL*8		SUMCAL	REAL*8
	WRITE5	REAL*8		WRITE5	REAL*8
TQHSBC	HST3D	REAL*8	TQSPRO	HST3D	REAL*8
	APLYBC	REAL*8		SUMCAL	REAL*8
	DUMP	REAL*8		WELLSS	REAL*8
	INIT2	REAL*8		WRITE5	REAL*8
	READ1	REAL*8			
4	SUMCAL	REAL#8	TQSSBC	HST3D	REAL*8
	WRITE5	REAL*8		APLYBC	REAL*8
		e e e e		DUMP	REAL*8
TQSAIF	HSTJD	REAL*8		INIT2	REAL*8
-	APLYBC	REAL*8		READ1	REAL*8
	DUMP	REAL#8		SUMCAL	REAL*8
	init2	REAL*8		WRITE5	REAL*8
	READ1	REAL*8			
	SUNCAL	REAL*8	TRVIS	HST3D	REAL*8
	WRITE5	REAL*8	•	DUMP	REAL*8
				etom1	REAL*8
TQSFBC	HST3D	REAL*8		READ1	REAL*8
	APLYBC	REAL*8		READ2	REAL*8
	DUMP	REAL*8		VISCOS	REAL*8
	INIT2	REAL*8		VSINIT	REAL*8
	READ1	REAL*8		WRITE2	REAL*8
	SUMCAL	REAL*8			
	WRITE5	REAL*8			

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Table 11.2--Cross-reference list of variables--Continued

Variable	Refer- encing	Variable	Variable name	Refer- encing	Variable
name	programs	type	II diuc	programs	type
TSBC	ASEMBL	REAL*8 DIMENSION(*)	TSYM	ASEMBL	REAL*8
	INIT3	REAL+8 DIMENSION(*)			
	SUHCAL	REAL*8 DIMENSION(*)	TSYP	ASEHBL	REAL*8
	WRITE3	REAL*8 DIMENSION (*)			
		•••	TSYX	COEFF	REAL*8 DIMENSION(*)
TSLBL	PLOTOC	CHARACTER#50		CRSDSP	REAL*8 DIMENSION(*)
				WRITE4	REAL+8 DIMENSION(*)
TSHAX	PLOTOC	REAL*8		•	
-			TSYZ	COEFF	REAL*8 DIMENSION(*)
TSMIN	PLOTOC	REAL*8		CRSDSP	REAL*8 DIMENSION(*)
				WRITE4	REAL*8 DIMENSION(*)
TSRES	HST3D	REAL*8			
	SUHCAL	REAL*8	TSZ	COEFF	REAL*8 DIMENSION(*)
	WRITE5	REAL*8		WRITE4	REAL*8 DIMENSION(*)
					5711 A0
TSRESF	HST3D	REAL+8	tszm	ASEMBL	REAL+8
	SUMCAL	REAL*8	-	4.000/01	551 × 40
	WRITE5	REAL*8	TSZP	ASEMBL	REAL*8
TSX	Coeff	REAL*8 DIMENSION(*)	TSZX	COEFF	REAL*8 DIMENSION(*)
	Write4	REAL*8 DIMENSION(*)		CRSDSP	REAL*8 DIMENSION(*)
	•			WRITE4	REAL*8 DIMENSION(*)
TSXM	ASEMBL	REAL*8			
			TSZY	COEFF	REAL*8 DIMENSION(*)
TSXP	ASEMBL	REAL*8		CRSDSP	REAL*8 DIMENSION(*)
				WRITE4	REAL*8 DIMENSION(*)
TSXY	COEFF	REAL*8 DIMENSION(*)			
	CRSDSP	REAL*8 DIMENSION(*)	TVD	HST3D	REAL*8 DIMENSION(10)
	WRITE4	REAL*8 DIMENSION(*)		DUMP	REAL*8 DIMENSION(10)
				etom1	REAL*8 DIMENSION(10)
TSXZ	COEFF	REAL*8 DIMENSION(*)		INIT2	REAL*8 DIMENSION(10)
	CRSDSP	REAL*8 DIMENSION(*)		READ1	REAL*8 DIMENSION(10)
	WRITE4	REAL*8 DIMENSION(*)		READ2	REAL*8 DIMENSION(10)
				WRITE2	REAL*8 DIMENSION(10)
TSY	COEFF	REAL*8 DIMENSION(*)			
	WRITE4	REAL*8 DIMENSION(*)			

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
TVFO	HST3D /	REAL*8 DIMENSION(10)	TWMAX	PLOTOC	REAL*8
	DUMP	REAL*8 DIMENSION(10)			
	ETOM1	REAL*8 DIMENSION(10)	TWMIN	PLOTOC	REAL*8
	INIT2	REAL*8 DIMENSION(10)	·		
	READ1	REAL*8 DIMENSION(10)	TWOPI	INIT2	REAL*8
	READ2	REAL*8 DIMENSION(10)	•		
	VISCOS	REAL*8 DIMENSION(10)	TWREND	HST3D	REAL*8
	WRITE2	REAL*8 DIMENSION(10)		WBBAL	REAL*8
				WELLSS	REAL*8
TVF1	HST3D	REAL*8 DIMENSION(10)		WELRIS	REAL*8
	DUMP	REAL*8 DIMENSION(10)			
	ETOM1	REAL*8 DIMENSION(10)	TWRK	WFDYDZ	REAL*8
	INIT2	REAL*8 DIMENSION(10)			
	READ1	REAL*8 DIMENSION(10)	TWSRKT	READ3	REAL*8 DIMENSION(*)
	READ2	REAL*8 DIMENSION(10)		WRITE3	REAL*8 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(10)			
•			TWSUR	WBBAL	REAL*8 DIMENSION(*)
TVZHC	HST3D	REAL*8 DIMENSION(5)		WELLSS	REAL*8 DIMENSION(*)
	DUMP	REAL*8 DIMENSION(5)		WRITES	REAL*8 DIMENSION(*)
	ETOM1	REAL*8 DIMENSION(5)			
	INIT2	REAL*8 DIMENSION(5)	TX	COEFF	REAL*8 DIMENSION(*)
	READ1	REAL*8 DIMENSION(5)	•	INIT2	REAL*8 DIMENSION(*)
	READ2	REAL*8 DIMENSION(5)		WRITE2	REAL*8 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(5)			
			TY	COEFF	REAL*8 DIMENSION(*)
TWKT	ASEMBL	REAL*8 DIMENSION(*)		INIT2	REAL*8 DIMENSION(*)
	WBBAL	REAL*8 DIMENSION(*)		WRITE2	REAL*8 DIMENSION(*)
	WELLSS	REAL*8 DIMENSION(*)			
	WRITE5	REAL*8 DIMENSION(*)	TZ	COEFF	REAL*8 DIMENSION(*)
			· · ·	IN1T2	REAL*8 DIMENSION(*)
TWLBL	PLOTOC	CHARACTER*50		WRITE2	REAL*8 DIMENSION(*)

Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
UO	BSODE	REAL*8	UBBLB	ETOM1	REAL*8 DIMENSION(*)
	INIT2	REAL*8		INIT2	REAL*8 DIMENSION(*)
	SUMCAL	REAL*8		READ2	REAL*8 DIMENSION(*)
Ul	BSODE	REAL*8	UBBRB	READ2	REAL*8
	COEFF	REAL#8			
	init2	REAL*8	UC	INIT2	REAL*8
	PLOT	INTEGER*4		PLOTOC	REAL*8
	SUMCAL	REAL*8		READ3	REAL*8
	WELLSS	REAL#8		SUHCAL	REAL*8
	WRITE2	REAL*8		WBBAL	REAL*8
	WRITE5	REAL+8		WELLSS	REAL*8
		•		WRITE3	CHARACTER*11
U2	COEFF	REAL*8			
	SUHCAL	REAL*8	UCBC	INIT3	REAL*8 DIMENSION(*)
	WRITE2	REAL*8		READ3	REAL*8 DIMENSION(*)
	WRITES	REAL#8			
			UCLM	WBBAL	REAL*8
U3	COEFF	REAL*8		Wellss	REAL*8
	WRITE2	REAL*8			
	WRITE5	REAL*8	UCLP	WBBAL	REAL*8
	•			WELLSS	REAL*8
U4	WRITE2	REAL*8			
	WRITE5	REAL*8	UCRBC	READ3	REAL*8
U5	WRITE2	REAL#8	UCROSC	ASEMBL	REAL*8
	WRITES	REAL*8		CRSDSP	REAL*8
U6	WRITE2	REAL*8	UCROST	ASEMBL	REAL*8
	WRITES	REAL*8		CRSDSP	REAL*8
				•	

UCWTCOEFFREAL*8UDXCOEFFREAL*8UDCINIT2REAL*8UDXDYINIT2REAL*8UDELYPLOTREAL*8UDXDYIINIT2REAL*8UDENCOEFFREAL*8UDXDYOINIT2REAL*8UDENCOEFFREAL*8UDXDZINIT2REAL*8UDENECETOM2REAL*8UDENSION(*)UDXYZINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDXYZINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*)UDYZINIT2REAL*8UDWKTITERREAL*8DIMENSION(*)UDYCOEFFREAL*8UDTINIT2REAL*8DIMENSION(*)UDYZINIT2REAL*8UDTINIT2REAL*8UEHWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLPWBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLPWBALREAL*8	Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable Lype
WELLSSREAL*8UDXDYIINIT2REAL*8UDELYPLOTREAL*8UDXDYOINIT2REAL*8UDENCOEFFREAL*8UDXDZINIT2REAL*8UDENCOEFFREAL*8UDXDZINIT2REAL*8UDENBCETOM2REAL*8UDXIXZINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDXYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDENLBINIT3REAL*8DIMENSION(*)UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*)UDYCOEFFREAL*8UDYWXTITERREAL*8DIMENSION(*)UDYCOEFFREAL*8UDTINIT2REAL*8DIMENSION(*)UDHWBBALREAL*8UDTINIT2REAL*8DIMENSION(*)UEHWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLMWBBALREAL*8	UCWT	COEFF	REAL*8	UDX	COEFF	REAL*8
UDELYPLOTREAL*8UDXDYIINIT2REAL*8UDENCOEFFREAL*8UDXDZINIT2REAL*8UDENCOEFFREAL*8UDXDZINIT2REAL*8UDENBCETOM2REAL*8DIMENSION(*) READ3UDXYZINIT2REAL*8UDENBCETOM2REAL*8DIMENSION(*) READ3UDXYZINIT2REAL*8UDENBCETOM2REAL*8DIMENSION(*) READ3UDXYZINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*) READ3UDYCOEFFREAL*8UDENLBETOM2REAL*8DIMENSION(*) READ3UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) READ3UDYCOEFFREAL*8UDYVKTITERREAL*8DIMENSION(*) REAL*8UDYCOEFFREAL*8UDTWKTITERREAL*8DIMENSION(*) REAL*8UEHWBBAL WELLSSREAL*8UDTHHCETOM1REAL*8DIMENSION(*) REAL*8UEHLMWBBAL WELLSSREAL*8	UDC	INIT2	REAL*8	UDXDY	INIT2	REAL*8
UDELYPLOTREAL*8UDXDYOINIT2REAL*8UDENCOEFF INIT2REAL*8UDXDZINIT2REAL*8UDENALREAL*8UDXDZINIT2REAL*8UDENBCETOM2 INIT3 REAL*8REAL*8UDXYZINIT2REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8DIMENSION(*) IDMENSION(*) READ3UDYZZINIT2REAL*8UDENLBETOM2 REAL*8REAL*8DIMENSION(*) INIT3 REAL*8UDYZZINIT2REAL*8UDENLBITTERREAL*8DIMENSION(*) READ3UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) REAL*8UDZCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) REAL*8UDZCOEFFREAL*8UDPWKTITERREAL*8DIMENSION(*) REAL*8UDEHWBBAL WELLSSREAL*8UDTHHCETOM1 INIT2REAL*8DIMENSION(*) REAL*8UEHLMWBBAL WBBAL REAL*8REAL*8		WELLSS	REAL*8			
UDENCOEFF INIT2REAL*8 REAL*8 SUMCAL WBBALREAL*8 REAL*8 DIMENSION(*) REAL*8UDXDZINIT2REAL*8UDENBCETOM2 INIT3 REAL*8REAL*8 DIMENSION(*) READ3UDXYZINIT2REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8 DIMENSION(*) READ3UDXYZINIT2REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8 DIMENSION(*) READ3UDXYZOINIT2REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8 DIMENSION(*) READ3UDYCOEFF REAL*8REAL*8UDNRBCREAD3REAL*8 DIMENSION(*) REAL*8UDZCOEFF REAL*8REAL*8UDYWKTITER WELLSS REAL*8REAL*8 REAL*8UDEH WBBAL WEBAL REAL*8REAL*8 REAL*8UDTHHCETOM1 INIT2 REAL*8REAL*8 DIMENSION(*) REAL*8WEHLP WBBAL WEBAL REAL*8REAL*8				UDXDYI	init2	REAL*8
UDENCOEFF INIT2 SUMCAL WBBALREAL*8 REAL*8UDXDZINIT2 INIT2REAL*8UDENBCETOM2 INIT3 REAL*8REAL*8DIMENSION(*) DIMENSION(*) READ3UDXYZINIT2 REAL*8REAL*8UDENBCETOM2 INIT3 REAL*8REAL*8DIMENSION(*) DIMENSION(*) READ3UDXYZINIT2 REAL*8REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8DIMENSION(*) DIMENSION(*) READ3UDXCOEFF REAL*8REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8DIMENSION(*) DIMENSION(*)UDYCOEFF REAL*8REAL*8UDNRBCREAD3REAL*8DIMENSION(*) REAL*8UDZCOEFF REAL*8REAL*8UDYWKTITERREAL*8UEH WBBAL WELLSSREAL*8REAL*8UDTINIT2 REAL*8REAL*8UEHLM WBBAL WELLSSREAL*8UDTHHCETOM1 INIT2REAL*8DIMENSION(*) REAL*8UEHLP WBBAL WEBAL REAL*8REAL*8	UDELY	PLOT	REAL*8			
INIT2REAL*8UDXDZINIT2REAL*8SUMCALREAL*8UDXYZINIT2REAL*8UDENBCETOM2REAL*8DIMENSION(*)UDXYZINIT2REAL*8UDENBCETOM2REAL*8DIMENSION(*)UDXYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDXYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDENLBREAD3REAL*8DIMENSION(*)UDYREAL*8REAL*8UDNRBCREAD3REAL*8DIMENSION(*)UDZINIT2REAL*8UDYKTITERREAL*8DIMENSION(*)UDZCOEFFREAL*8UDTINIT2REAL*8DIMENSION(*)UDYREAL*8REAL*8UDTINIT2REAL*8DIMENSION(*)UEHWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLMWBBALREAL*8UDTHHCINIT2REAL*8DIMENSION(*)UEHLPWBBALREAL*8			· · ·	UDXDYO	INIT2	REAL*8
SUMCAL WBBALREAL*8UDXYZINIT2REAL*8UDENBCETOM2 INIT3 READ3REAL*8DIMENSION(*) READ3UDXYZIINIT2REAL*8UDENLBETOM2 READ3REAL*8DIMENSION(*) READ3UDXYZOINIT2REAL*8UDENLBETOM2 INIT3 REAL*8REAL*8DIMENSION(*) REAL*8UDYCOEFFREAL*8UDENLBETOM2 READ3REAL*8DIMENSION(*) READ3UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) READ3UDZCOEFFREAL*8UDPWKTITERREAL*8UMENSION(*) REAL*8UDZCOEFFREAL*8UDTINIT2 REAL*8REAL*8UEHWBBAL WELLSSREAL*8UDTINIT2 REAL*8REAL*8UEHLMWBBAL WELLSSREAL*8UDTHHCETOM1 INIT2REAL*8DIMENSION(*) REAL*8UEHLPWBBAL WEBAL REAL*8	UDEN	COEFF	REAL*8			
WBBALREAL*8UDXYZINIT2REAL*8UDENBCETOM2REAL*8DIMENSION(*)UDXYZIINIT2REAL*8INIT3REAL*8DIMENSION(*)UDXYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*)UDYZZINIT2REAL*8UDYWKTITERREAL*8DIMENSION(*)UDHWBBALREAL*8UDTINIT2REAL*8DIMENSION(*)UEHWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLPWBBALREAL*8		INIT2	REAL*8	UDXDZ	INIT2	REAL*8
UDENBCETOM2 INIT3 REAL*8REAL*8DIMENSION(*) INIT3 REAL3UDXYZIINIT2REAL*8UDENLBETOM2 INIT3 REAL3REAL*8DIMENSION(*) INIT3 REAL*8UDYCOEFFREAL*8UDENLBETOM2 INIT3 REAL*8REAL*8DIMENSION(*) INIT3 REAL*8UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) INIT2UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) UDYUDYCOEFFREAL*8UDYITERREAL*8UDZCOEFFREAL*8UDTINIT2 WELLSSREAL*8UEHWBBAL WELLSSREAL*8UDTHHCETOM1 INIT2REAL*8DIMENSION(*) UEHLPWBBAL WBBAL REAL*8REAL*8		SUMCAL	REAL*8			
INIT3 READ3REAL*8DIMENSION(*) IMENSION(*)UDXYZOINIT2REAL*8UDENLBETOH2 INIT3 REAL*8REAL*8DIMENSION(*) IMENSION(*)UDYCOEFFREAL*8UDNRBCREAD3REAL*8DIMENSION(*) IMERSION(*)UDZINIT2REAL*8UDNRBCREAD3REAL*8DIMENSION(*) IMERSION(*)UDZCOEFFREAL*8UDPWKTITERREAL*8UDZCOEFFREAL*8UDTINIT2REAL*8UEHWBBAL WELLSSREAL*8UDTINIT2REAL*8DIMENSION(*) WELLSSUEHLMWBBAL WELLSSREAL*8UDTHHCETOM1 INIT2REAL*8DIMENSION(*) IMENSION(*)UEHLPWBBAL WBBAL REAL*8REAL*8		WBBAL	REAL*8	UDXYZ	INIT2	REAL*8
READ3REAL*8DIMENSION(*)UDXYZOINIT2REAL*8UDENLBETOM2REAL*8DIMENSION(*)UDYCOEFFREAL*8INIT3REAL*8DIMENSION(*)UDYDZINIT2REAL*8UDNRBCREAD3REAL*8DIMENSION(*)UDZCOEFFREAL*8UDPWKTITERREAL*8UDZCOEFFREAL*8UDTINIT2REAL*8UEHWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLPWBBALREAL*8	UDENBC	ETOM2	REAL*8 DIMENSION(*)	UDXYZI	INIT2	REAL*8
UDENLBETOM2 INIT3 REAL3REAL*8 DIMENSION(*) DIMENSION(*) READ3UDYCOEFFREAL*8UDNRBCREAD3REAL*8 DIMENSION(*) REAL*8UDTDZINIT2REAL*8UDPWKTITERREAL*8UDZCOEFFREAL*8UDTINIT2REAL*8UDZCOEFFREAL*8UDTINIT2REAL*8UEHWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTHHCETOM1REAL*8 DIMENSION(*) REAL*8 DIMENSION(*)UEHLPWBBALREAL*8		INIT3	REAL*8 DIMENSION(*)		•	
INIT3 READ3REAL*8DIMENSION(*) READ3UDYDZINIT2REAL*8UDNRBCREAD3REAL*8UDZCOEFFREAL*8UDPWKTITERREAL*8UEHWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLPWBBALREAL*8		READ3	REAL*8 DIMENSION(*)	UDXYZO	INIT2	REAL*8
READ3REAL*8DIMENSION(*)UDYDZINIT2REAL*8UDNRBCREAD3REAL*8UDZCOEFFREAL*8UDPWKTITERREAL*8UEHWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTHHCETOM1REAL*8DIMENSION(*)UEHLPWBBALREAL*8	UDENLB	ETOM2	REAL*8 DIMENSION(*)	UDY	COEFF	REAL*8
UDNRBCREAD3REAL*8UDZCOEFFREAL*8UDPWKTITERREAL*8UEHWBBALREAL*8UDTINIT2REAL*8UEHLMWBBALREAL*8UDTHHCETOM1REAL*8 DIMENSION(*)UEHLPWBBALREAL*8		INIT3	REAL*8 DIMENSION(*)			*
UDPWKT ITER REAL*8 UDT INIT2 REAL*8 WELLSS REAL*8 UDT ETOM1 REAL*8 DIMENSION(*) INIT2 REAL*8 DIMENSION(*) UEHLP WBBAL REAL*8		READ3	REAL*8 DIMENSION(*)	UDYDZ	INIT2	REAL*8
UDT INIT2 REAL*8 WELLSS REAL*8 UDTHHC ETOM1 REAL*8 DIMENSION(*) INIT2 REAL*8 DIMENSION(*) UEHLP WBBAL REAL*8	UDNRBC	READ3	REAL*8	UDZ	COEFF	REAL*8
UDT INIT2 REAL*8 WELLSS REAL*8 UDTHHC ETOM1 REAL*8 DIMENSION(*) INIT2 REAL*8 DIMENSION(*) UEHLP WBBAL REAL*8		ITER	REAL*8	UEH	WBBAL	REAL*8
WELLSS     REAL*8     UEHLM     WBBAL     REAL*8       UDTHHC     ETOM1     REAL*8     DIMENSION(*)     WELLSS     REAL*8       INIT2     REAL*8     DIMENSION(*)     UEHLP     WBBAL     REAL*8					WELLSS	REAL*8
UDTHHC ETOM1 REAL*8 DIMENSION(*) INIT2 REAL*8 DIMENSION(*) UEHLP WBBAL REAL*8	UDT		•	,		
UDTHHC ETOM1 REAL*8 DIMENSION(*) INIT2 REAL*8 DIMENSION(*) UEHLP WBBAL REAL*8		WELLSS	REAL*8	UEHLM	WBBAL	
INIT2 REAL*8 DIMENSION(*) UEHLP WBBAL REAL*8					WELLSS	REAL*8
	UDTHHC		•••			
READ2 REAL*8 DIMENSION(*)				UEHLP	WBBAL	REAL*8
WRITE2 REAL*8 DIMENSION(*) UFRAC COEFF REAL*8		WRITE2	REAL*8 DIMENSION(*)	UFRAC	COEFF	REAL*8
UDTIM COEFF REAL*8 UFX COEFF REAL*8	UDTIM	COEFF	REAL*8	UFX	COEFF	REAL*8

Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
UFY	COEFF	REAL+8	UNIGRZ	HST3D	LOGICAL*4
				INIT2	LOGICAL*4
UFZ	COEFF	REAL*8		READ2	LOGICAL*4
				WRITE2	LOGICAL*4
UGDELX	INIT2	REAL=8			
			UNITEP	REWI3	CHARACTER*10
UCDELY	INIT2	REAL*8	·	WRITE3	CHARACTER*10
UGDELZ	INIT2	REAL#8	UNITH	HST3D	CHARACTER*3
				DUMP	CHARACTER*3
UHRBC	READ 3	REAL*8		INITI	CHARACTER*3
				PLOTOC	CHARACTER*3
UHWT	COEFF	REAL*8		READ1	CHARACTER*3
				REWI3	CHARACTER*3
ພະ	READ2	INTEGER*4		WRITE2	CHARACTER*3
				WRITE3	CHARACTER*3
UKHCBC	ETOM1	REAL*8 DIMENSION(*)		WRITE4	CHARACTER*3
	init2	REAL*8 DIMENSION(*)		WRITE5	CHARACTER*3
	READ2	REAL*8 DIMENSION(*)			
	•		Unithf	HST3D	CHARACTER*7
UKLB	eton1	REAL*8 DIMENSION(*)		DUNP	CHARACTER*7
	init2	REAL*8 DIMENSION(*)		INITI	CHARACTER*7
	READ2	REAL*8 DIMENSION(*)		PLOTOC	CHARACTER#7
				READI	CHARACTER*7
UKRB	READ2	REAL*8		REWI 3	CHARACTER*7
				WRITE2	CHARACTER*7
UNICRX	hst3d	LOGICAL*4		WRITE3	CHARACTER*7
	INIT2	LOGICAL*4		WRITE4	CHARACTER*7
	READ2	LOGICAL#4		WRITE5	CHARACTER*7
	WRITE2	LOGICAL#4			
UNICRY	hst3d	LOGICAL*4			
	INIT2	LOGICAL*4			
	READ2	LOGICAL*4		1	
	WRITE2	LOGICAL*4	(		

Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
UNITL	HST3D	CHARACTER*2	UNITT	HST3D	CHARACTER*1
	DUMP	CHARACTER*2		DUMP	CHARACTER*1
	INITI	CHARACTER*2		INITI	CHARACTER*1
	PLOTOC	CHARACTER*2		PLOTOC	CHARACTER*1
	READI	CHARACTER*2		READI	CHARACTER*1
1.	REWI 3	CHARACTER*2		REWI3	CHARACTER*1
	WRITE2	CHARACTER*2		WRITE2	CHARACTER*1
k	WRITE3	CHARACTER*2		WRITE3	CHARACTER*1
$\lambda = -\ell$	WRITE4	CHARACTER*2		WRITE4	CHARACTER*1
	WRITE5	CHARACTER*2		WRITE5	CHARACTER*1
NITM	HST3D	CHARACTER*2	UNITTM	HST3D	CHARACTER*3
	DUMP	CHARACTER*2		DUMP	CHARACTER*3
	INITI	CHARACTER*2		INIT1	CHARACTER*3
	PLOTOC	CHARACTER*2		PLOTOC	CHARACTER*3
	READI	CHARACTER*2		READ1	CHARACTER*3
	REWI 3	CHARACTER*2		REWI3	CHARACTER*3
	WRITE2	CHARACTER*2		WRITE2	CHARACTER*3
	WRITE3	CHARACTER*2		WRITE3	CHARACTER*3
	WRITE4	CHARACTER*2		WRITE4	CHARACTER*3
	WRITE5	CHARACTER*2		WRITE5	CHARACTER*3
NITP	HST3D	CHARACTER*3	UNITVS	REWI 3	CHARACTER*9
	DUMP	CHARACTER*3		WRITE2	CHARACTER*9
	INITI	CHARACTER*3		WRITE3	CHARACTER*9
,	PLOTOC	CHARACTER*3			_
	READ1	CHARACTER*3	UNLP	PLOT	INTEGER*4
	REWI 3	CHARACTER*3			
	WRITE2	CHARACTER*3	UP	WELLSS	REAL*8
	WRITE3	CHARACTER*3			
	WRITE4	CHARACTER*3	UP 1	READ3	REAL*8
	WRITE5	CHARACTER*3		WRITE3	CHARACTER*11

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Table 11.2--Cross-reference list of variables--Continued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
UP2	READ3	BEAL+8	UQHLP	WBBAL	REAL*8
	WRITE3	CHARACTER*11		WELLSS	REAL*8
UPABD	INIT2	REAL*8	uqs	INIT3	REAL*8
UPHILB	ETOM2	REAL*8 DIMENSION(*)	UQW	WELLSS	REAL*8
	INIT3	REAL*8 DIMENSION(*)			
	READ3	REAL*8 DIMENSION(*)	UQWLM	WBBAL	REAL*8
				WELLSS	REAL*8
UPHIM	APLYBC	REAL*8		tinn (*	554540
		5711 + 40	UQWLP	WBBAL	REAL*8
UPS	PLOTOC	REAL*8		Wellss	REAL*8
UPTC	COEFF	REAL*8	UQWLYV	WBBAL	REAL*8
UPW	PLOTOC	REAL#8	UQWM	WBBAL	REAL*8
			•	WELLSS	REAL*8
UPWKT	ITER	REAL*8			
	WELLSS	REAL*8	UQWV	WBBAL	REAL*8
				WELLSS	REAL*8
UPWKTS	WELLSS	REAL*8			
			UR1	ASEMBL	REAL*8
UQ	INIT3	REAL*8			
•	READ3	REAL*8	UR2	ASEMBL	REAL*8
	WRITE3	CHARACTER*11			
			URH	ASEMBL	REAL*8
UQCLM	WBBAL	REAL*8			
-	WELLSS	REAL*8	URS	ASEMBL	REAL*8
UQCLP	WBBAL	REAL*8	UT	INIT2	REAL*8
	WELLSS	REAL*8		PLOTOC	REAL*8
				READ3	REAL*8
UQH	INI <b>T</b> 3	REAL*8		SUMCAL	REAL*8
,				VSINIT	REAL*8
uqhlm	WBBAL	REAL*8		WRITE3	CHARACTER*11
	WELLSS	REAL*8			
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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
UTBC	ETOM2	REAL*8 DIMENSION(*)	UVAR	READ2	REAL*8
	INIT3	REAL*8 DIMENSION(*)			
	READ3	REAL*8 DIMENSION(*)	UVEL.	COEFF	REAL*8
UTLM	WBBAL	REAL*8	UVFLM	WELLSS	REAL*8
	WELLSS	REAL*8		· .	
			UVFLP	WELLSS	REAL*8
UTLP	WELLSS	REAL*8			~~~
			UVIS	COEFF	REAL*8
UTRBC	READ3	REAL*8			
	D1 0700		UVISLB	ETOM2	REAL*8 DIMENSION(*)
UTS	PLOTOC	REAL#8		INIT3	REAL*8 DIMENSION(*)
UTTC	COEFF	REAL*8	•	READ3	REAL*8 DIMENSION(*)
	51.0500		UVSRBC	READ3	REAL*8
UTW	PLOTOC	REAL*8			
UTXH	ASEMBL	REAL*8	UVWLM	WELLSS	REAL*8
U VVII	NOCUUL	a.cm,, "G	UVWLP	WELLSS	REAL#8
UTXP	ASEMBL	REAL#8	UYWLE	MEPP99	NPUR.A
VAAL	<b>43 111 24 111 24</b>	presi 50 V	UWI	INIT2	REAL*8
UTYM	ASEMBL	REAL*8	UWA	****	
<b>1 8 8 8 8</b>			UXX1	SOR2L	RFAL*8
UTTP	ASEMBL	REAL*8	UARE	*****	· · · · · · · · · · · · · · · · · · ·
			UXX2	SOR2L	REAL*8
UTZH	ASEMBL	REAL*8		÷ = = - =	
			UYMAX	PLOT	REAL*8
UTZP	ASEMBL	REAL*8			
		. · · · · · · · · · · · · · · · · · · ·	UYMIN	PLOT	REAL*8
UVAIFC	etom1	REAL*8 DIMENSION(*)			
	INIT2	REAL*8 DIMENSION(*)	UZELB	ERROR2	REAL*8 DIMENSION(*)
	READ2	REAL*8 DIMENSION(*)		etom1	REAL*8 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(*)		INIT2	REAL*8 DIMENSION(*)
			1	READ2	REAL*8 DIMENSION(*)

Variable	Refer- encing		riable	Variable	Refer- encing		riable
name	programs	i.	type	name	programs	1	type
UZERB	READ2	REAL*8		VISCTR	HST3D		DIMENSION(10)
					DUMP	REAL*8	DIMENSION(10)
VA	ASEMBL	REAL*8	DIMENSION(7,*)		etom1	REAL*8	DIMENSION(10)
	D4DES	REAL*8	DIMENSION(7,*)		1NIT2	REAL*8	DIMENSION (10)
	ITER	REAL*8	DIMENSION(7,*)		READ1	REAL*8	DIMENSION (10)
	SOR2L		DIMENSION(7,*)		READ2	REAL*8	DIMENSION (10)
					VISCOS	REAL*8	DIMENSION(10)
VAIFC	APLYBC	REAL*8	DIMENSION(*)		WRITE2		DIMENSION (10)
	INIT2		DIMENSION(*)				
			•••	VISLBC	APLYBC	REAL*8	DIMENSION(*)
VAR	REWI	REAL*8			INIT3	REAL*8	DIMENSION(*)
	REWI 3	REAL*8	DIMENSION(3)		WRITE3	REAL*8	DIMENSION(*)
VASBC	ASEMBL	REAL*8	DIMENSION(7,*)	VISOAR	HST3D	REAL*8	
	SBCFLO	REAL*8	DIMENSION(7,*)		APLYBC	REAL*8	
					DUMP	REAL*8	
VAW	ASEMBL	REAL*8	DIMENSION(7,*)		ETOH1	REAL*8	
	WBCPLO	REAL*8	DIMENSION(7,*)		INIT2	REAL*8	
			• • •		READ1	REAL*8	
VDATA	VSINIT	REAL*8	DIMENSION(*)		READ2	REAL*8	
				•	WRITE2	REAL*8	
VELWRK	WFDYDZ	REAL*8					
		-		VISTFO	HST3D	REAL*8 1	DIMENSION (10)
VIS	APLYBC	REAL*8	DIMENSION(*)		DUHP		DIMENSION (10)
	COEFF		DIMENSION(*)		ETOMI		DIMENSION (10)
	INIT2		DIMENSION(*)		INIT2		DIMENSION (10)
	SUHCAL		DIMENSION(*)		READ1		DIMENSION (10)
	WELLSS	REAL*8	DIMENSION(*)		READ2		DIMENSION (10)
	WRITE2	REAL*8	DIMENSION(*)		VISCOS		DIMENSION (10)
	WRITES		DIMENSION(*)		WRITE2		DIMENSION(10)

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Variable	Refer- encing	Variable	Variable	Refer- encing	Variable
name	programs	type	name	programs	type
VISTP1	HST3D	REAL*8 DIMENSION(10)	WO	HST3D	REAL*8
	DUMP	REAL*8 DIMENSION(10)		DUMP	REAL*8
	ETOM1	REAL*8 DIMENSION(10)		INIT2	REAL*8
	INIT2	REAL*8 DIMENSION(10)		INIT3	REAL*8
	READI	REAL*8 DIMENSION(10)		READ1	REAL*8
	READ2	REAL*8 DIMENSION(10)		READ2	REAL*8
	WRITE2	REAL*8 DIMENSION(10)	ι.	READ3	REAL*8
				SUMCAL	REAL*8
VPA	RST3D	REAL*8 DIMENSION(*)	Y	VISCOS	REAL*8
	ASEMBL	REAL*8 DIMENSION(*)		WBBAL	REAL*8
	DUMP	REAL*8 DIMENSION(*)		WELLSS	REAL*8
	INITI	REAL*8 DIMENSION(*)		WRITE2	REAL*8
	INIT2	REAL*8 DIMENSION(*)		WRITE3	REAL*8
	ITER	REAL*8 DIMENSION(*)		WRITE5	REAL*8
	L2SOR	REAL*8 DIMENSION(*)			1
	READI	REAL*8 DIMENSION(*)	W1	HST3D	REAL*8
1	SUMCAL	REAL*8 DIMENSION(*)		DUMP	REAL*8
	WBBAL	REAL*8 DIMENSION(*)		INIT2	REAL*8
	WELLSS	REAL*8 DIMENSION(*)		INIT3	REAL*8
				READ1	REAL#8
VREF	VSINIT	REAL*8		READ2	REAL*8
	•	•		READ3	REAL*8
VSTLOG	VSINIT	REAL*8 DIMENSION(16)		VISCOS	REAL*8
		· · ·		WRITE2	REAL*8
VXX	COEFF	REAL*8 DIMENSION(*)		WRITE3	REAL*8
	WRITE4	REAL#8 DIMENSION(*)		WRITE5	REAL*8
VII	COEFF	REAL*8 DIMENSION(*)	WBOD	ETOM1	REAL*8 DIMENSION(*)
	WRITE4	REAL*8 DIMENSION(*)		INIT2 READ2	REAL*8 DIMENSION(*) REAL*8 DIMENSION(*)
VZZ	COEFF	REAL*8 DIMENSION(*)		WELRIS	REAL*8
	WRITE4	REAL*8 DIMENSION(*)		WRITE2	REAL*8 DIMENSION(*)

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-338 -63 -63 Table 11.2--Cross-reference list of variables--Continued

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	Refer-			Refer-	
Variable	encing	Variable	Variable	encing	Variable
name	programs	type	name	programs	type
WCAIF	APLYBC	REAL*8 DIMENSION(*)	WIDLBL	PLOT	CHARACTER*80
	INIT2	REAL*8 DIMENSION(*)		PLOTOC	CHARACTER*80
	SUMCAL	REAL*8 DIMENSION(*)			
			WQMETH	ASEMBL	INTEGER*4 DIMENSION(*)
NCF	EKROR2	REAL*8 DIMENSION(*)		ERROR2	INTEGER*4 DIMENSION(*)
	INIT2	REAL*8 DIMENSION(*)		ERROR3	INTEGER*4 DIMENSION(*)
	READ2	REAL*8 DIMENSION(*)		ITER	INTEGER*4 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(*)		PLOTOC	INTEGER*4
				READ2	INTEGER*4 DIMENSION(*)
WCLBL1	WRITE2	CHARACTER*60 DIMENSION(0:	5)	READ3	INTEGER*4 DIMENSION(*)
				SUHCAL	INTEGER*4 DIMENSION (*)
WCLBL2	WRITE2	CHARACTER*50 DIMENSION(0:	2)	WBBAL	INTEGER*4 DIMENSION(*)
				WELLSS	INTEGER*4 DIMENSION(*)
WFICUM	INIT2	REAL*8 DIMENSION(*)		WRITE2	INTEGER*4 DIMENSION(*)
	SUHCAL	REAL*8 DIMENSION(*)		WRITE3	INTEGER*4 DIMENSION(*)
	WELLSS	REAL*8 DIMENSION(*)		WRITE5	INTEGER*4 DIMENSION(*)
	WRITES	REAL*8 DIHENSION(*)			
			WRANGL	READ2	REAL*8 DIMENSION(*)
WFPCUM	INIT2	REAL*8 DIMENSION(*)		WELRIS	REAL*8
	SUMCAL	REAL*8 DIMENSION(*)		WRITE2	REAL*8 DIMENSION(*)
	Wellss	REAL*8 DIMENSION(*)			
	WRITE5	REAL*8 DIMENSION(*)	WRCALC	HST3D	LOGICAL*4
				DUMP	LOGICAL*4
WHICUM	INIT2	REAL*8 DIMENSION(*)		etom1	LOGICAL*4
	SUHCAL	REAL*8 DIMENSION(*)		INIT2	LOGICAL*4
	WELLSS	REAL*8 DIMENSION(*)		READ1	LOGICAL*4
	WRITES	REAL*8 DIMENSION(*)		<b>KEAD2</b>	LOGICAL*4
				WEBAL	LOGICAL*4
WHPCUM	INIT2	REAL*8 DIMENSION(*)		WELLSS	LOGICAL*4
	SUHCAL	REAL*8 DIMENSION(*)		WELRIS	LOG1CAL*4
	WELLSS	REAL*8 DIMENSION(*)		WRITE2	LOGICAL#4
	WRITE5	REAL*8 DIMENSION(*)		WRITE5	LOGICAL*4
WI	INIT2	REAL*8 DIMENSION(*)			
	WELLSS	REAL*8 DIMENSION(*)			
	WRITE2	REAL*8 DIMENSION(*)	<i>r</i>		

Variable	Refer-	Variable	Variable	Refer- encing programs	Variable
name	programs	type	name		type
WRID	ETOMI	REAL*8 DIMENSION(*)	WIMY	ASEMBL	REAL*8
	READ2	REAL*8 DIMENSION(*)			
	WELRIS	REAL*8	WTMZ	ASEMBL	REAL*8
	WRITE2	REAL*8 DIMENSION(*)			
			WTPX	ASEMBL	REAL*8
WRIDT	HST3D	REAL*8			
	WELRIS	REAL*8	WTPY	ASEMBL	REAL*8
	WFDYDZ	REAL*8			
		···· - · ·	WTPZ	ASEMBL	REAL*8
WRISL	ETOMI	REAL*8 DIMENSION(*)			
	READ2	REAL*8 DIMENSION(*)	x	COEFF	REAL*8 DIMENSION(*)
	WELLSS	REAL*8 DIMENSION(*)		ERROR2	REAL*8 DIMENSION(*)
	WELRIS	REAL*8		ETOM1	REAL*8 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(*)		INIT2	REAL*8 DIMENSION(*)
				MAP2D	REAL*8 DIMENSION(*)
WRRUF	ETOM1	REAL*8 DIMENSION(*)		PLOT	REAL*8
·	READ2	REAL*8 DIMENSION(*)		READ2	REAL*8 DIMENSION(*)
	WELRIS	REAL*8		WRITE2	REAL*8 DIMENSION(*)
	WRITE2	REAL*8 DIMENSION(*)		WRITE5	REAL*8 DIMENSION(*)
				ZONPLT	REAL#8 DIMENSION(*)
WSICUM	INIT2	REAL*8 DIMENSION(*)			•
	SUMCAL	REAL*8 DIMENSION(*)	XO	INIT2	REAL*8
	WELLSS	REAL*8 DIMENSION(*)			•
	WRITE5	REAL*8 DIMENSION(*)	<b>X1</b>	MAP2D	REAL*8
· .			•	ZONPLT	REAL*8
WSPCUM	INIT2	REAL*8 DIMENSION(*)			
	SUMCAL	REAL*8 DIMENSION(*)	X2	MAP2D	REAL*8
	WELLSS	REAL*8 DIMENSION(*)	••••	ZONPLT	REAL*8
	WRITES	REAL*8 DIMENSION(*)			
	**************************************	titere o perimipant(")	XARG	INTERP	REAL*8
WT	COEFF	REAL*8			
** #	UVHL E	алан 140 °° 67	XC	PLOT	REAL*8 DIMENSION(*)
WTHX	ASEMBL	REAL*8		ZONPLT	REAL*8

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Table 11.2--Cross-reference list of variables--Continued

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/ariable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable Lype
XD.	ZONPLT	REAL#8	Y	COEFF	REAL*8 DIMENSION(*)
				ETOM1	REAL*8 DIMENSION(*)
KI	WELRIS	REAL*8	•	INIT2	REAL*8 DIMENSION(*)
				HAP2D	REAL*8 DIMENSION(*)
XINC	PLOT	REAL+8		PLOT	REAL*8 DIMENSION(6)
				READ2	REAL*8 DIMENSION(*)
XLBL	PLOT	CHARACTER*30		WRITE2	REAL*8 DIMENSION(*)
				WRITE5	REAL*8 DIMENSION(*)
XMAX	PLOT	REAL*8		ZONPLT	REAL*8 DIMENSION(*)
	ZONPLT	REAL+8			
			YO	INIT2	REAL*8
OHIN	PLOT	REAL*8			
	ZONPLT	REAL*8	Y1	MAP2D	REAL*8
				WELLSS	REAL#8
KO	PLOT	REAL*8 DIMENSION(*)		WFDYDZ	REAL*8
				ZONPLT	REAL*8
(PRNT	PLOT	REAL#8			
		•	¥2	MAP2D	REAL*8
KS	INTERP	REAL*8 DIMENSION(*)		ZONPLT	REAL*8
XTOT	MAP2D	REAL*8	YARG	INTERP	REAL+8
xx	INIT2	REAL+8	YC	PLOT	REAL*8 DIMENSION(*)
-	SOR2L	REAL*8 DIMENSION(*)		ZONPLT	REAL*8
	TOPEP	REAL*8			
	VISCOS	REAL*8	YD	ZONPLT	REAL*8
	VSINIT	REAL*8			
			YLBL	PLOT	CHARACTER*40
KXN	SOR2L	REAL*8 DIMENSION(*)		-	
			YHAX	PLOT	REAL*8
XYRAT	ZONPLT	REAL*8		ZONPLT	REAL*8

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Table 11.2Cross-reference	list of	variablesContinued

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Variable name	Refer- encing programs	Variable type	Variable name	Refer- encing programs	Variable type
YMAXHV	BSODE	REAL*8 DIMENSION(2,12)	YYMXSV	BSODE	REAL*8 DIMENSION(2)
YMIN	PLOT ZONPLT	REAL*8 Real*8	YYN	BSODE	REAL*8 DIMENSION(2)
			YYNMI	BSODE	REAL*8 DIMENSION(2)
VIIIV	BSODE	REAL*8 DIMENSION(2,12)	•••••		
			YYSAVE	BSODE	REAL*8 DIMENSION(2)
YNM1HV	BSODE	REAL*8 DIMENSION(2,12)			
			<b>Z</b> .	APLYBC	REAL*8 DIMENSION(*)
<b>YO</b>	PLOT	REAL*8 DIMENSION(*)		ASEMBL	REAL*8 DIMENSION(*)
10	WELLSS	REAL*8		BSODE	REAL*8
	WFDYDZ	REAL*8		COEFF	REAL*8 DIMENSION(*)
				ERROR2	REAL*8 DIMENSION(*)
YPOSUP	HST3D	LOGICAL*4		ETOM1	REAL*8 DIMENSION(*)
	READ3	LOGICAL*4		INIT2	REAL*8 DIMENSION(*)
	WRITE3	LOGICAL*4		READ2	REAL*8 DIMENSION(*)
	WRITE5	LOGICAL*4		SUMCAL	REAL*8 DIMENSION(*)
				WELLSS	REAL*8 DIMENSION(*)
YREF	VSINIT	REAL*8		WRITE2	REAL*8 DIMENSION(*)
				WRITE5	REAL*8 DIMENSION(*)
YS	INTERP	REAL*8 DIMENSION(*)			
			<b>ZO</b>	INIT2	REAL*8
YTOT	MAP2D	REAL*8			
	· · · · · ·	<b>**</b>	Z1	MAP2D	REAL <sup>+8</sup>
YY	BSODE	REAL*8 DIMENSION(2)		· · ·	
·	VSINIT	REAL*8	Z2	MAP2D	REAL+8
	WELRIS	REAL*8 DIMENSION(2)			
	WFDYDZ	REAL*8 DIMENSION(2)	ZA	BSODE	REAL*8
YYERR	BSODE	REAL*8 DIMENSION(2)	<b>2CHARS</b>	MAP2D	CHARACTER*1 DIMENSION(0:31)
	WELRIS	REAL*8 DIMENSION(2)			
			ZEBRA	MAP2D	LOGICAL*4
YYMAX	BSODE	REAL*8 DIMENSION(2)		WRITE5	LOGICAL*4
	WELRIS	REAL*8 DIMENSION(2)			

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Table 11.2--Cross-reference list of variables--Continued

Variable name	Refer- encing programs		riable Lype	Variable name	Refer- encing programs	v	ariable type
ZELBC	APLYBC	REAL*8	DIMENSION(*)	ZT	HST3D	REAL*8	DIMENSION (10)
	ASEMBL		DIMENSION(*)		DUHP	REAL*8	DIMENSION(10)
	ERROR3	REAL#8	DIMENSION(*)		etom1	REAL*8	DIMENSION(10)
	INIT2	REAL+8	DIMENSION(*)		INIT2	REAL*8	DIMENSION(10)
	SUHCAL	REAL+8 1	DIMENSION(*)		READ1	REAL*8	DIMENSION(10)
	WRITE2	REAL*8	DIMENSION(*)		READ2	REAL*8	DIMENSION(10)
					WRITE2	REAL*8	DIMENSION(10)
zhcbc	APLYBC	REAL*8	Dimension(*)				
	etom1	REAL*8 1	dimension(*)	ZTHC	HST3D	REAL*8	DIMENSION(5)
	INIT2	REAL*8	dimension(*)		DUMP	REAL*8	DIMENSION(S)
	READ2	REAL*8	DIMENSION(*)		etom1	REAL*8	DIMENSION(5)
	WRITE2	REAL+8	dimension(*)		INIT2	REAL*8	DIMENSION(5)
		•			READ	REAL*8	DIMENSION(5)
ZPINIT	HST3D	REAL*8			READ2	REAL*8	DIMENSION(5)
	DUMP	REAL#8			WRITE2	REAL*8	DIMENSION(5)
	etom1	REAL*8					-
	init2	REAL*8		ZU	BSODE	REAL*8	
	READI	REAL*8					
	READ2	REAL*8		ZWK	WELRIS	REAL*8	
	WRITE2	REAL*8			WFDYDZ	REAL*8	
ZPOSUP	HST3D	LOGICAL	*4	ZWKT	WELLSS	REAL*8	
	READ3	LOGICAL	k4				
	WRITE3	LOGICAL	≜4				

Table 11.2--Cross-reference list of variables--Continued

WRITES

LOGICAL\*4

## 11.3 CROSS-REFERENCE LIST OF COMMON BLOCKS

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A cross-reference list of common blocks (table 11.3) shows in which subprograms each named common block appears. Blank common is used only for the two variably-partitioned arrays. Generally, the common block names relate to the subprogram in which the variables of that common block are defined first. All common blocks are contained in the main program.

1. A. A. A.

Софиол		Common		Common		Common	
block	Referencing	block name	Referencing programs	block name	Referencing programs	block name	Referencing programs
name	programs						
ASA	MAIN	IN1B	MAIN	INIG	MAIN	INIM	HAIN
	ASEMBL	,	DUMP		DUMP		DUHP
	CRSDSP	·	ERROR 1		INITI		INITI
			INITI		ITER		ITER
CCC	MAIN		READ1		READ 1		READI
	ASEMBL		WRITEI				SUMCAL
	CALC			INIH	HAIN		
		INIC	HAIN		DUMP	INIP	HAIN
ERI	HAIN		DUMP		INITI		DUMP
	ERROR 1		INITI		L2SOR		INITI
	ERROR2		L2SOR		READ1		READI
	ERROR3		READI				
	ERRPRT			1111	MAIN	1N1Q	HAIN
	INIT2	INID	MAIN		DUMP	•	DUMP
	IREWI		DUMP		INITI		INITI
	1 TER		INITI	,	READ 1		ITER
	READI		ITER				READI
4	READ2		READ1	INIJ	HAIN		
	READ3				ASEMBL	INIS	MAIN
	REWI	INIE	MAIN		DUHP		DUMP
	REWI 3		ASEMBL		INITI		INIT1
	SOR2L		DUHP		READ1		PLOTOC
	SUHCAL		INITI				READI
			READI	INIL	MAIN		WRITE2
INIA	MAIN				LUMP		WRITE3
	DUMP	INIF	MAIN		INITI		WRITE4
	INITI		DUMP		INIT2		WRITES
	READ 1		INITI		READ1		
			INIT2		WBBAL		
			READ1		WELLSS		

Table 11.3--Cross-reference list of common blocks

Common	· ·····	Common	•	Common		Common	
block	Referencing	block	Referencing	block	Referencing	block	Referencing
name	programs	name	programs	name	programs	name	programs
INIT	MAIN	IN2BV	MAIN	IN2F	MAIN	IN2L	MAIN
	DUMP		BLKDAT		APLYBC		CLOSE
	INIT1		DUMP		DUMP		DUMP
	PLOTOC		INIT2		INIT2		INIT2
	READI		READ1		READ1		READI
	REWI 3		TOFEP				
	WRITE2			IN2H	MAIN	ITA	MAIN
	WRITE3	IN2C	MAIN		APLYBC		ASEMBL
	WRITE4	·	APLYBC		CALC		ITER
	WRITES		ASEMBL		CLOSE		WRITE5
			DUMP		COEFF		
1N2A	MAIN		INIT2		DUMP	L2SAV	MAIN
	APLYBC		READI		ERROR3		DUMP
	ASEMBL		SUMCAL		ETOM2		L2SOR
	CALC		WBBAL		INIT2		READI
	COEFF		WELLSS		INIT3		WRITE5
	DUMP		WRITE2		ITER		· ·
	ERROR3		WRITES		READ1	ORA	MAIN
	INIT2				READ3		ASEMBL
	INIT3	IN2D	MAIN		SUMCAL		D4DES
	READI		DUMP		WBBAL		DUMP
	READ3		INIT2		WELLSS		INITI
	SUMCAL		READI		WRITE2		ORDER
	WELLSS		VISCOS		WRITE3		READ1
	WELRIS		,		WRITE4		SBCFLO
	•	IN2E	MAIN		WRITES		WRITEL
IN2B	MAIN		CALC	•			
	BLKDAT		DUMP	IN2I	MAIN		
	DUMP		INIT2		APLYBC		
	INIT2		ITER		DIMP		
	READI		<b>READ1</b>		INIT2		
	TOFEP		SUMCAL		· READ1		
			WBBAL		SUMCAL		
			WELLSS	1	WRITE2		
			WFDYDZ		WRITE5		

Table 11.3--Cross-reference list of common blocks--Continued

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Connon		Common		Common		Common	
block name	Referencing programs	block name	Referencing programs	block name	Referencing programs	block name	Referencing programs
RDIA	HAIN	RD1B	HAIN	RD1D	HAIN	RDIE	HAIN
	APLYBC		READ1		ASEHBL		APLYBC
	ASEHBL		WRITE1		CLOSE		ASEMBL
	CALC				COEFF		DUMP
	COEFF	RD1C	HAIN		DUHP		ERRORI
	CRSDSP		APLYBC		ERROR2		ERROR2
	DUMP		ASEHBL		ERROR3		ERROR3
	ERROR2		COEFF		ETOH 1		ETOHI
	ERROR3		CRSDSP	·	INITI		ETOH2
	ETOMI		D4DES		INIT2		INITI
	ETOM2		DUHP		INIT3		INIT3
	INITI		ERROR 1		IREWI		INIT3
	INIT2		ERROR2		ITER		ITER
	INIT3		ETOM1		L2SOR		READ1
	ITER		ETOH2		PLOTOC		READ2
	PLOTOC		INITI		READI		READ3
	READ1		INIT2		READ2		SBCFLO
	READ2		INIT3		READ3		SUHCAL
	READ3		IREWI		REWI		WRITE1
	SUHCAL		ITER		REWI 3		WRITE2
	VISCOS		ORDER		SUNCAL		WR1TE3
	WBBAL		PLOTOC		<b>WBBAL</b>		WRITE5
	WELLSS		PRNTAR		WELLSS		
	WRITE1		READ1		WRITE1	RD1F	MAIN
	WRITE2		READ2		WRITE2		COEFF
	WRITE3		READ3		WRITE3		DUHP
	WRITE4		REWI		WRITE4		ERROR 1
	WRITE5		REWI3		WRITES		ERROR2
			SBĊFLO		ZONPLT		ETOH1
DIAC	MAIN		SOR2L				INITI
	DUMP		SUMCAL				INIT2
	READI		WBBAL				READI
	WRITEI		WELLSS				READ2
			WRITEL				WRITE1
			WRITE2		•		WRITE2
			WRITE3				ZCNPLT
			WRITE4		1		
1			WRITE5	1			
(				1			

## Table 11.3--Cross-reference list of common blocks--Continued

Common		Common		Common	·····	Common	
block name	Referencing programs	block	Referencing programs	block na <del>me</del>	Referencing programs	block name	Referencing programs
RDIG	MAIN	RD2B	MAIN	RD2E	MAIN	RD2G	
10010	APLYBC	NDED	APLYBC	ND2E	ASEMBL	RDZG	MAIN DUMP
	DUMP		ASEMBL		COEFF		ERROR2
	ERROR 1	,	CALC		DUMP		ETOM1
	ERROR2		COEFF		ETOHI		INIT2
	ETOHI		CRSDSP		INIT2		READI
1	INITI		DUMP		INIT3	. 1	READ2
	INIT2		ETOMI		PLOTOC		VISCOS
	READI	•	INIT2		READI		WRITE2
	READ2		READ1		READ2		
	WRITE1		READ2		READ3	RD2GV	MAIN
	WRITE2		SUMCAL		SUMCAL		DUMP
		· · ·	WELLSS		VISCOS		ETOMI
RDIH	MAIN		WELRIS		WBBAL		INIT2
	ASEMBL		WFDYDZ		WELLSS		READI
	COEFF		WRITE2		WRITE2		READ2
	DUMP		WRITES		WRITE3		VISCOS
	ERROR 1				WRITES		WRITE2
	INIT1	RD2C	MAIN				
	INIT2		DUMP	RD2F	MAIN	RD2II	MAIN
	ITER		ETOM1		DUMP		APLYBC
	READ1		INIT2		ETOM1		DUMP
	READ2		READI		INIT2		ERROR2
	SBCFLO		READ2		READ1		ETOM1
	WRITE1		WRITE2		READ2		INIT2
	WRITE2				WRITE2		READ1
	WRITE4	RD2D	MAIN				READ2
	WRITE5		INIT2				WRITE2
			READ2				
RD2A	MAIN		WRITE2			RD21V	MAIN
•	DUMP						APLYBC
	ERROR2						BLKDAT
	INIT2						DUMP
	READ1						READI
	READ2				1		READ2
	WRITE2						WRITE2

Table 11.3--Cross-reference list of common blocks--Continued

Common		Common		Common		Common	
block	Referencing	block	Referencing	block	Referencing	block	Referencing
nane	programs	name	programs	name	programs	name	programs
RD2J	MAIN	RD2L	MAIN	RD2P	MAIN	RD2U	MAIN
	APLYBC		DUMP		DUMP		DUMP
	ASEHBL		INIT2		ETOH 1		IREWI
	DUHP	•	ITER		INIT2		READI
	ETOHI		READ1		READ 1		READ2
	INIT2		READ2		READ2		REWI
	READ1		WRITE2		WELLSS		REWI3
	READ2				WRITE2		SUHCAL
	SUHCAL	RD2M	HÁIN				
	TOFEP		CLOSE	RD2Q	HAIN	RD2V	MAIN
	WBBAL		DUHP		DUMP		DUMP
	WELLSS		READ1		INIT2		INIT2
	WELRIS		READ2		ITER		READ1
	WFDYDZ		READ3		READ 1		READ2
	WRITE2		WELLSS		READ2		READ3
	WRITE5		WELRIS		WRITE2		SUMCAL
			WRITE2				WRITE2
RD2K	MAIN		WR1TE3				WRITE3
	DUHP		WRITES	RD2S	MAIN		WRITE4
	ERROR2				APLYBC		WR1TES
	ETOM1	RD2N	MAIN		ASEMBL		
	INIT2		DUMP		COEFF		
	READI		ETOH1		DUHP	RD2W	ETOH1
	READ2		INIT2		READ1		
	WRITE2		ITER		READ2	RD3A	HAIN
			READI		SBCFLO		ERROR3
RD2KV	MAIN		READ2		SUHCAL		ETOM2
	DUMP		WBBAL		WELLSS		INIT3
	ETOM		WELLSS		WRITE2		READ3
	INIT2		WELRIS		WRITES		WRITE3
	READ1		WRITE2				
	READ2		WRITES	RD2T	MAIN	RD3B	MAIN
	WRITE2				DUHP		COEFF
					INIT2		ERROR3
					ITER		ETOH2
					READ1		INIT3
					READ2		<b>READ</b> 3
					SOR2L		WRITE3
,					WRITE2		
ſ				(			

Table 11.3--Cross-reference list of common blocks--Continued

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Common		Common		Common		Common	
block	Referencing	block	Referencing	block	Referencing	block	Referencing
name	programs	name	programs	name	programs	name	programs
RD3C	MAIN	RD31	MAIN	SCA	MAIN	SORA	MAIN
	READ3		CLOSE		COEFF		SOR2L
	WRITE4		COEFF		ITER		WRITE5
	WRITE5		DUMP	•	SUMCAL		
			ERROR3		WRITES	WRA	MAIN
RD3D	MAIN		ETOM2				WELRIS
	ERROR3		INIT3	SCB	MAIN		WFDYDZ
	READ3		READ3		APLYBC		
	WRITE3		SUMCAL		SUMCAL	WRB	MAIN
	WRITES		WRITE1		WRITE5		BSODE
			WRITE3				WELRIS
RD3E	MAIN		WRITE4	SCC	MAIN		
	CLOSE		WRITE5		SUMCAL	WRC	MAIN
	DUMP .				WELLSS		WELRIS
	READ3				WRITE5		WFDYDZ
<b>RD3EV</b>	MAIN	-				WSSA	MAIN
	ERROR3						WBBAL
	ETOM2						WELLSS
	INIT3						WELRIS
	READ3						WFDYDZ
	WRITE3						
	WRITES.						
RD3F	MAIN	х Х					
= =	READ3						
RD3G	MAIN						
	COEFF		,				,
	READ3						
	WRITE4						

Table 11.3--Cross-reference list of common blocks--Continued

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11.4 COMMENT FORM AND MAILING-LIST REQUEST

If you wish to be placed on a mailing list for revisions of the program documentation and announcements of new releases of this program, please return this page to:

> HST3D Code Custodian U.S. Geological Survey Water Resources Division Box 25046, Mail Stop 413 Denver Federal Center Denver, Colorado 80225

Any comments on the program or documentation will be appreciated.

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Telephone Number ( )