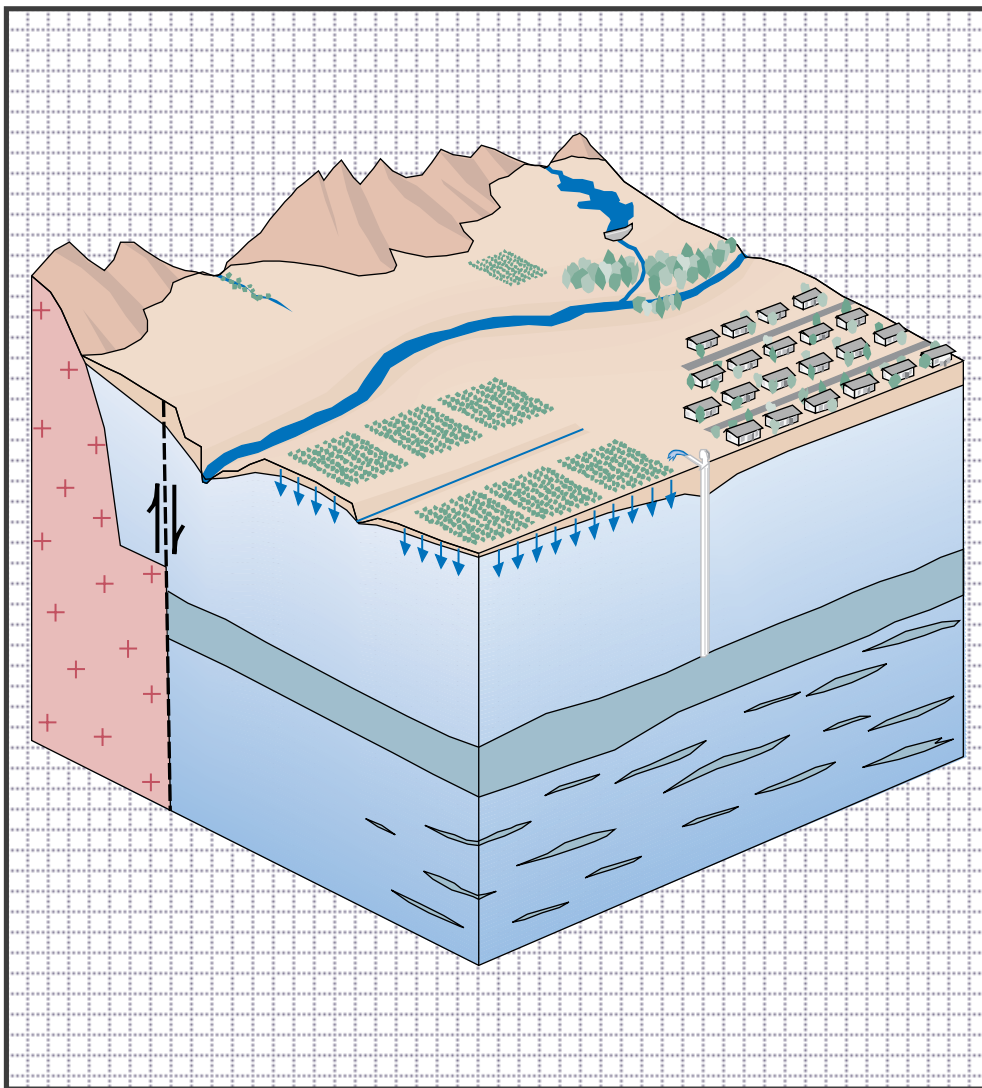


MODFLOW-2000, THE U.S. GEOLOGICAL SURVEY MODULAR GROUND-WATER MODEL—USER GUIDE TO MODULARIZATION CONCEPTS AND THE GROUND-WATER FLOW PROCESS

Open-File Report 00-92



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PREFACE

This report describes an enhanced version of the U.S. Geological Survey modular ground-water model, called MODFLOW-2000, for which the structure has been expanded to facilitate the solution of multiple related equations. The performance of the program has been tested in a variety of applications. Future applications, however, might reveal errors that were not detected in the test simulations. Users are requested to notify the U.S. Geological Survey of any errors found in this User Guide or the computer program by using the address on the back of the report title page. Updates might occasionally be made to both the User Guide and to MODFLOW-2000. Users can check for updates on the Internet at URL http://water.usgs.gov/software/ground_water.html/.

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MODFLOW-2000, The U.S. Geological Survey Modular Ground-Water Model—User Guide to Modularization Concepts and the Ground-Water Flow Process

By Arlen W. Harbaugh, Edward R. Banta, Mary C. Hill, and Michael G. McDonald

ABSTRACT

MODFLOW is a computer program that numerically solves the three-dimensional ground-water flow equation for a porous medium by using a finite-difference method. Although MODFLOW was designed to be easily enhanced, the design was oriented toward additions to the ground-water flow equation. Frequently there is a need to solve additional equations; for example, transport equations and equations for estimating parameter values that produce the closest match between model-calculated heads and flows and measured values. This report documents a new version of MODFLOW, called MODFLOW-2000, which is designed to accommodate the solution of equations in addition to the ground-water flow equation. This report is a user's manual. It contains an overview of the old and added design concepts, documents one new package, and contains input instructions for using the model to solve the ground-water flow equation.

INTRODUCTION

MODFLOW is a computer program that simulates three-dimensional ground-water flow through a porous medium by using a finite-difference method (McDonald and Harbaugh, 1988). MODFLOW was designed to have a modular structure that facilitates two primary objectives: ease of understanding and ease of enhancing. Ease of understanding was an objective because U.S. Geological Survey technical managers generally believe that modelers should understand how a model works in order to use it properly. Ease of enhancement was an objective because experience showed that there was a continuing need for new capabilities. Examples of enhancements are reports by Prudic (1989), Hill (1990), Leake and Prudic (1991), Goode and Appel (1992), Harbaugh (1992), McDonald and others (1992), Hsieh and Freckleton (1993), Leake and others (1994), Fenske and others (1996), and Leake and Lilly (1997).

MODFLOW was originally documented by McDonald and Harbaugh (1984). As with most computer programs that are used over a long time period, MODFLOW underwent several overall updates. The second version of MODFLOW is documented in McDonald and Harbaugh (1988), and this version is often called MODFLOW-88 to distinguish it from other versions. A third version is called MODFLOW-96 (Harbaugh and McDonald, 1996a and 1996b).

In addition to the enhancements and updates, the U.S. Geological Survey developed two major extensions to MODFLOW—MODFLOWP (Hill, 1992) and MOC3D (Konikow and others, 1996). MODFLOWP and MOC3D solve equations in addition to the ground-water flow equation. MODFLOWP solves a MODFLOW calibration problem by calculating values of selected input data that result in the best match between measured and model calculated values, and MOC3D solves the solute-transport equation for concentration.

Although MODFLOW was originally designed to facilitate change, its design concepts did not include solving equations other than the ground-water flow equation. As a result, incorporating capabilities such as those added through MODFLOWP and MOC3D was not as straightforward from the programmer's or user's

perspective as were the other enhancements that dealt only with the ground-water flow equation. Therefore, MODFLOW-2000 has been developed to facilitate the addition of multiple types of equations. Ease of understanding continues to be included as an objective of the design. There is also an objective to minimize changes that would impact existing MODFLOW users.

For many data input quantities, MODFLOW-2000 allows definition using parameter values, each of which can be applied to data input for many grid cells. In combination with new multiplication and zone array capabilities, the parameters make it much easier to modify data input values for large parts of a model. Defined parameters also can have associated sensitivities calculated and be modified to attain the closest possible fit to measured hydraulic heads, flows, and advective travel. This is accomplished using the Observation, Sensitivity, and Parameter-Estimation Processes of MODFLOW-2000, which are documented by Hill and others (2000).

This report first documents the enhanced modular structure that is incorporated within MODFLOW-2000 to accommodate the solution of multiple equations. An overview of the new design concepts is provided from the perspective of a user; the detailed design information needed by programmers will be provided in separate documentation as the information becomes available. This report also documents the incorporation of the ground-water flow equation within the new structure, including the documentation of a new package called the Layer-Property Flow Package and input instructions for using MODFLOW-2000 to simulate ground-water flow. A user will also need to refer to earlier MODFLOW reports, such as McDonald and Harbaugh (1988), to find information about how the ground-water flow equation is solved by using the finite-difference method. The parts of MODFLOW-2000 that solve additional equations also are documented in separate reports, such as Konikow and others (1996) and Hill and others (2000).

Although some new data-input capabilities have been added to MODFLOW-2000, most of the input data used by earlier versions of MODFLOW to simulate ground-water flow can be read without change. A data conversion program is provided to convert the parts of the input data that require change.

DESIGN CONCEPTS

This section of the report discusses several general aspects of MODFLOW-2000. The first involves the old and new modularization concepts and how they work together; these are presented in the first three sections below. Next, MODFLOW-2000 allows system characteristics to be defined by using parameters, and the basic ideas behind this change are discussed. Finally, a basic change in the major output files is discussed.

Previously Existing Design Concepts—Packages, Procedures, and Modules

MODFLOW-2000 expands upon the modularization approach that was originally included in MODFLOW (McDonald and Harbaugh, 1988). The design of MODFLOW can be viewed from two perspectives. The first perspective is the user's perspective. A user views the program according to its capability to simulate different kinds of ground-water flow problems. To facilitate this perspective, the program is divided into pieces called packages. Each hydrologic capability, such as leakage to rivers, recharge, and evapotranspiration, that is included within the ground-water flow equation is a separate package. Further, because there are many methods for solving the simultaneous equations resulting from the finite-difference method, each solution method is a package. The Preconditioned Conjugate Gradient (PCG2 of Hill, 1990) and Strongly Implicit Procedure (SIP) are examples of solution methods implemented as packages in MODFLOW. Finally, the remainder of the code that is not included in a hydrologic or solution package becomes the Basic (BAS) Package. BAS provides overall program control.

The second perspective is the perspective of the programmer, for whom the goal is to break the program into pieces that make the program logic as simple as possible while maintaining the desired functionality. In MODFLOW, these pieces are called procedures. The program logic, which is typically represented by a flowchart, is thus represented by describing the sequence of procedures. Examples of procedures are the Read and Prepare Procedure in which user-specified data are read and prepared for use and the Formulate Procedure in which the finite-difference equation is constructed for each model cell.

To make it possible to divide easily the program into both packages and procedures as desired, the program code is broken into smaller pieces called modules. Each module contains the program code within a single procedure for a single package. The code for a package can therefore be created by collecting all of the modules that correspond to the package. Likewise, the code for any procedure can be created by collecting all of the modules that correspond to the procedure. The MAIN program creates the required procedural view of the program by invoking the various modules in the proper procedural sequence. New packages can thus be added to MODFLOW by providing the new modules and modifying the MAIN program to invoke the new modules in the proper procedural sequence.

In summary, there are three modularization entities in all the earlier versions of MODFLOW: procedures, packages, and modules. The module is the most fundamental entity. Modules can be grouped according to packages or procedures in order to obtain the user's or programmer's perspective, respectively.

Added Design Concept—Processes

To incorporate the solution of multiple related equations, a fourth modularization entity called a process is added. A process is a part of the code that solves a fundamental equation by a specified numerical method. For example, the solution of the ground-water flow equation using the finite-difference method, which is the original MODFLOW, is now called the Ground-Water Flow (GWF) Process. The GWF Process includes all aspects of solving the flow equation, including the formulation of the finite-difference equations, data input, solving the resulting simultaneous equations, and data output. If a different method, the finite-element method for example, were used, then this would become a separate process. Note that the solution of the set of simultaneous equations within the GWF Process is not a process. Rather, this is only one part of the broader process of solving the ground-water flow equation by using the finite-difference method. Accordingly, as previously mentioned, there are several solver packages that can be used to solve the simultaneous equations, and these are all considered to be part of the GWF Process.

The Ground-Water Transport (GWT) Process (formerly referred to as MOC3D) is an example of another process. The GWT Process solves the solute-transport equation as described by Konikow and others (1996) by using the numerical method known as the method of characteristics. As with the ground-water flow equation, there are many numerical methods that can be used to solve the solute-transport equation. A fundamentally different method would be implemented as a separate process. Variants of the method of characteristics could be incorporated as part of the GWT Process. Model calibration as implemented by Hill and others (2000) requires three processes in addition to the GWF Process. The Observation (OBS) Process calculates simulated values that are to be compared to measurements, calculates the sum of squared, weighted differences between model values and observations, and calculates sensitivities related to the observations. The Sensitivity (SEN) Process solves the sensitivity equation for hydraulic heads throughout the grid, and the Parameter-Estimation (PES) Process solves the modified Gauss-Newton equation to minimize an objective function to find optimal parameter values.

All processes are divided into packages, procedures, and modules in the same way as in previous versions of MODFLOW. Therefore, in MODFLOW-2000 a module has three levels of identification— process, package, and procedure. The procedures within a process must be defined as required to program that process efficiently, and there need be no commonality of procedures among processes. When procedures of two or more processes are similar, however, there is some benefit from choosing the same procedure name. The naming convention selected for MODFLOW-2000 requires that packages that are represented in multiple processes use the same package name. For example, the modules in the Sensitivity Process that deal with river seepage using the River (RIV) Package of the Ground-Water Flow Process must be designated as the RIV Package of the Sensitivity Process.

It is assumed that every non-Ground-Water Flow Process within MODFLOW-2000 operates with the Ground-Water Flow Process. There is no assumption, however, that all the non-Ground-Water Flow processes work together. For example, the Parameter-Estimation Process by Hill and others (2000) does not work with the GWT Process. The documentation for each process should define the other processes with which it is compatible.

In addition to processes that solve equations, MODFLOW-2000 has the Global Process (GLO), which controls overall program operation and sets up data structures that can be used by all processes. Some of the initialization parts of the original MODFLOW have been put into the Global Process so that the Ground-Water

Flow Process is somewhat different in detail from the original MODFLOW. This is indicated in the MODFLOW-2000 flowchart presented in the next section.

Overall Structure

Depending on what processes are included along with the Ground-Water Flow Process, MODFLOW's structure can be significantly more complex than it was originally. For example, when the Observation, Sensitivity, and Parameter-Estimation Processes are added, a new parameter estimation loop is added to the structure, which allows the Ground-Water Flow Process to be executed multiple times, and a new parameter-sensitivity loop is added. A simplified flowchart of the Global, Ground-Water Flow, Observation, Sensitivity, and Parameter-Estimation Processes combined is shown in figure 1. To illustrate the changes compared to the original, the procedures and program loops of the Global and Ground-Water Flow Processes, which comprise the functionality of the original MODFLOW, are identified with asterisks. Normally, each procedure would be shown as a separate box; however, some of the sequential procedures are combined into a single box to save space in the illustration. Further, figure 1 excludes some of the less important procedures of the Observation, Sensitivity, and Parameter-Estimation Processes to minimize the size and complexity of the illustration. Enough of the structure is shown to illustrate the expansion that is typical when using multiple processes.

Most of the procedures listed in figure 1 are described by McDonald and Harbaugh (1988). A new procedure required by the Parameter-Estimation Process is Rewind (RW). The Rewind Procedure is needed to allow the ground-water flow equation to be solved multiple times without totally redefining the way that the Ground-Water Flow Process reads and stores data. This procedure rewinds the input files to their beginning before restarting the Ground-Water Flow Process. Thus, the input files are reread each time the flow equation is solved.

Parameters and Variables

Another change in MODFLOW-2000 is the addition of parameters as used in parameter-estimation terminology. In earlier MODFLOW documentation, the term parameter was generally used to mean any input data value. For example, the recharge flux at a cell is a parameter by this definition. However, a more restricted definition of "parameter" is commonly used when dealing with statistical parameter-estimation theory (Draper and Smith, 1998; Ott, 1993). To make it possible to smoothly incorporate parameter estimation into MODFLOW-2000 and provide an additional, convenient approach for specifying input data, the more restricted meaning is used in this report. Thus, here a parameter is a single value that is given a name and determines the value of a variable used in the finite-difference ground-water flow equation at one or more model cells. The definition of a parameter specifies which variable is being defined and the cells for which the parameter applies. For example, a parameter might define the aquifer hydraulic conductivity for a group of cells in a model layer, or a parameter might define the riverbed conductance for one or more reaches of a river. Parameters are defined as part of the Ground-Water Flow Process of MODFLOW-2000, and they can be used by other processes. In addition to facilitating the incorporation of the Observation, Sensitivity, and Parameter-Estimation Processes, the parameter approach for model input can make data input easier. For a full explanation of how to define parameters, refer to the Using Parameters in the Ground-Water Flow Process Section of this report.

In order to be consistent with this new usage of the term "parameter" in MODFLOW-2000, the term "auxiliary parameter" as used in MODFLOW-96 (Harbaugh and McDonald, 1996a, p. 26) has been changed to "auxiliary variable."

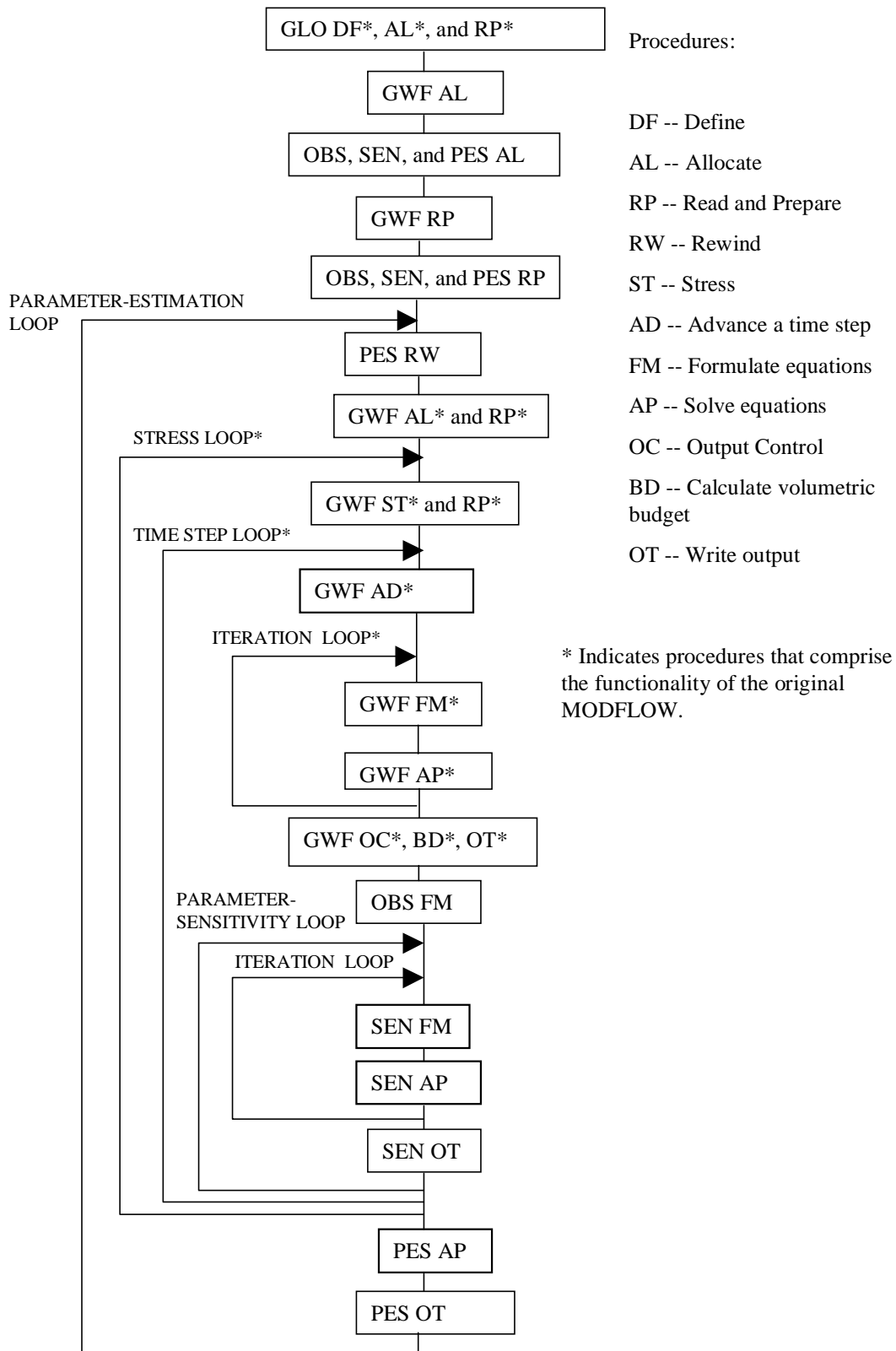


Figure 1. -- Flowchart of Global (GLO), Ground-Water Flow (GWF), Observation (OBS), Sensitivity (SEN), and Parameter Estimation (PES) Processes.

Listing Output

Earlier versions of MODFLOW always produced a file called the listing file. With the addition of the Parameter-Estimation Process, in which the parameter-estimation loop (fig. 1) may be repeated several times during the estimation of parameters, the potential for excessively long output files arises. During each parameter-estimation iteration, the Ground-Water Flow Process executes and produces written output related to one simulation of the ground-water flow problem, and the Sensitivity Process executes and produces written output related to each parameter considered. If all this output is written to a single file, then that file will increase in size, possibly dramatically, with each iteration of the Parameter-Estimation Process. In most cases, during parameter estimation, results of intermediate simulations are of little or no interest, and the output from these simulations is unneeded.

To provide users with the option to retain only the output of most interest and limit the amount of output generated, MODFLOW-2000 can produce either one or two listing output files. When the user specifies that one output file is to be generated, all listing output from the model is written to that file. When the user specifies that two output files are to be generated, output that applies to the overall model run is written to a file called the GLOBAL file, but most of the output from Ground-Water Flow and Sensitivity Processes is written to a file called the LIST file. Further, when a new iteration begins, the LIST file is erased and generated anew. As a result, most of the output from the Ground-Water Flow and Sensitivity Processes is limited to that produced during the most recently executed parameter-estimation iteration. Thus, the use of two files avoids generating excessively large amounts of output.

If the Parameter-Estimation Process is not being used, the amount of output is the same when either one or two output files are used because no process is executed more than once. However, two files can still be used in this situation to divide the output into two pieces. This is a matter of user preference. If the Parameter-Estimation Process is used, two output files are highly recommended. For additional information see Hill and others (2000).

As described in the Input Instructions Section of this report, the use of one or two listing files is determined by which file types are defined in the name file. File types LIST and GLOBAL indicate that the respective listing files are to be used (file type, as used for MODFLOW, is defined in the Activating Capabilities and Opening Files Using the Name File Section of this report). If only one of these file types is specified in the name file, then all the listing output is written to that file.

The rest of this section summarizes the contents of the GLOBAL and LIST files, with the assumption that the user has specified that both GLOBAL and LIST files be created. The GLOBAL file contains information that applies to the model run as a whole. Output from the Define (DF), Allocate (AL), and Read and Prepare (RP) procedures of the Global, Observation, Sensitivity, and Parameter-Estimation Processes (fig. 1) is written to the GLOBAL file. The output includes the names and types of all files that are opened. Input for the Global Process is echoed to the GLOBAL file; this information includes a description of the spatial and temporal discretization to be used. If the Observation, Sensitivity, or Parameter-Estimation Processes are being used, their input is echoed. Input for the selected solver package is echoed. The GLOBAL file also shows the parameter definitions read from the input files for the individual packages that are used to simulate various boundary conditions and stresses for the Ground-Water Flow Process.

If the Observation Process is active, the GLOBAL file also contains information that is calculated during the model run. If the Observation Process is active while the Sensitivity and Parameter-Estimation Processes are inactive, summary information related to the fit of the model calculations to the observed values is written to the GLOBAL file. If, in addition to the Observation Process, the Sensitivity Process is active, the GLOBAL file also includes information that quantifies the sensitivity of the model-calculated equivalents of observations to the parameter values. If, in addition to the Observation and Sensitivity Processes, the Parameter-Estimation Process is active, the GLOBAL file also lists information related to parameter estimation. For additional information about the functioning of MODFLOW-2000 when the Observation, Sensitivity, and Parameter-Estimation Processes are active, see Hill and others (2000; for example, table 3).

The LIST file contains output from the procedures of the Ground-Water Flow Process that appear within the parameter-estimation loop in figure 1. This output includes allocation information, values used by the Ground-Water Flow Process for layers and individual cells in solving the flow equation, and such calculated model results

as hydraulic head, drawdown, and volumetric budget. When the Observation Process is active, the LIST file also contains model-calculated equivalents to the observations and related statistics.

The Example Section of this report includes output from model runs that illustrate the use of GLOBAL and LIST files in a ground-water flow simulation. One model run uses a single LIST file, whereas the second uses both LIST and GLOBAL files.

GLOBAL PROCESS

As stated in the Design Concepts Section, the Global Process has been included in MODFLOW-2000 to control overall program flow, open files, and read global data such as space and time discretization. Unlike other processes, the Global Process does not actually solve an equation.

Activating Capabilities and Opening Files Using the Name File

MODFLOW-2000 activates capabilities and opens files the same way that MODFLOW-96 does. There is a single name file that contains the names of most of the files used by the Ground-Water Flow, Observation, Sensitivity, and Parameter-Estimation Processes. The name file also includes file types that determine which program options are activated. For example, if the name file contains a line beginning with file type RIV, then the River Package is activated. Instructions for preparing the name file are included in the Input Instructions Section of this report. Example name files are included in the Examples Section of this report.

Space Discretization

The Global Process reads information that defines the physical size of the finite-difference grid. As in all earlier MODFLOW versions, the finite-difference grid in MODFLOW-2000 is assumed to be rectangular horizontally, while the grid can be distorted vertically (fig. 2). However, unlike previous versions of MODFLOW, MODFLOW-2000 always requires the definition of the complete geometry of the each cell. Previous versions of MODFLOW do not require direct definition of vertical cell geometry for confined layers because the cell geometry is already incorporated within transmissivity (hydraulic conductivity of a cell times cell thickness), storage coefficient (specific storage times cell thickness), and vertical leakance (vertical hydraulic conductivity divided by the vertical distance between two nodes). By not directly specifying vertical cell geometry for confined layers, the amount of input data and computer memory requirements are minimized. MODFLOW-2000 always defines vertical cell geometry, however, because other processes, such as transport modeling, may need this information even if the Ground-Water Flow Process does not. Further, the need to minimize the use of computer memory is not as great as when MODFLOW was first developed.

The horizontal grid dimensions are specified in variables DELR and DELC (fig. 2A). Columns are numbered starting from the left side of the grid. Rows are numbered starting from the upper edge (plan view) of the grid. $DELR_j$ is the width of the cells (from the left side to the right side) in column j . That is, all the cells in a column have the same width, and there is one value of DELR for each of the NCOL columns in the model grid. Similarly, $DELC_i$ is the width of cells (from the top to the bottom in plan view) in row i , and there is one value of DELC for each of the NROW rows in the model grid.

Layers are numbered starting from the top layer and going down (fig. 2B). Elevation of the top of layer 1 is defined in addition to the bottom elevation of every layer. The elevation information can be used to calculate the thickness of all cells. Below each layer except the bottom layer, there can also be a confining bed through which only vertical flow is simulated. Simulating confining beds by this method often is called the Quasi-Three-Dimensional (Quasi-3D) Approach (McDonald and Harbaugh, 1988, p. 5-18). There is no requirement to use the Quasi-3D Approach; that is, any confining bed can be simulated using one or more distinct model layers as desired. Further, there is no requirement that all processes support the use of the Quasi-3D Approach. The GWT Process, for example, does not support the calculation of concentrations in confining beds in which the Quasi-3D

Approach is used (Konikow and others, 1996, p. 31). For these confining beds, the elevation of the bottom of the bed is also defined. The elevation information can be used to calculate the thickness of the confining beds.

As shown in figure 2B, the user must specify the top elevation of layer 1 and the bottom elevations of all model layers and confining beds. The top elevation of other model layers and confining beds need not be separately specified because the top elevation is the same as the bottom elevation of the layer or confining bed that is immediately above. There is no requirement to use distorted layers as shown in the illustration. Flat layers can be used by simply specifying a constant value for the elevation of each layer.

The space discretization information is included in a new file called the discretization file. Previously, this information was included in the Block-Centered Flow Package file.

Time Discretization

Time discretization in MODFLOW-2000 is dealt with much as in earlier versions of MODFLOW. The fundamental component of time discretization is the time step. Time steps are grouped into stress periods. Time dependent input data can be changed every stress period.

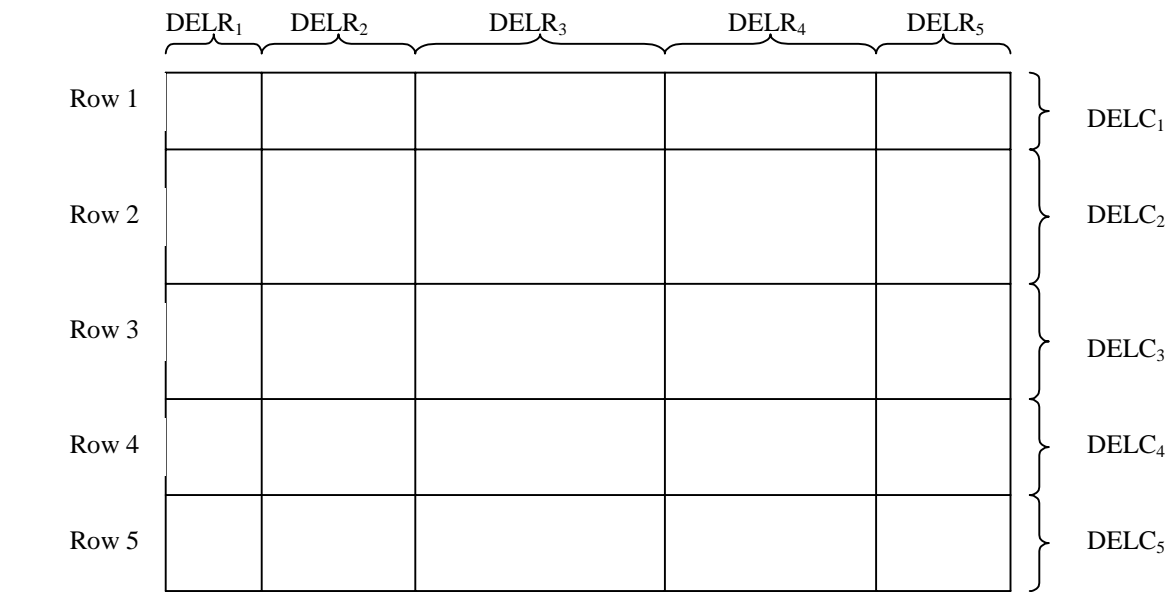
For each stress period, the user specifies the total length (PERLEN), the number of time steps (NSTP), and the multiplier for the length of successive time steps (TSMULT). That is, the length of time step n is the length of time step $n-1$ times TSMULT. A series of numbers in which each successive value is a constant times the previous value is called a geometric series. The length of the first time (Δt_1) step can be determined from the following equation for a geometric series:

$$\Delta t_1 = \text{PERLEN} \left(\frac{\text{TSMULT} - 1}{\text{TSMULT}^{\text{NSTP} - 1}} \right).$$

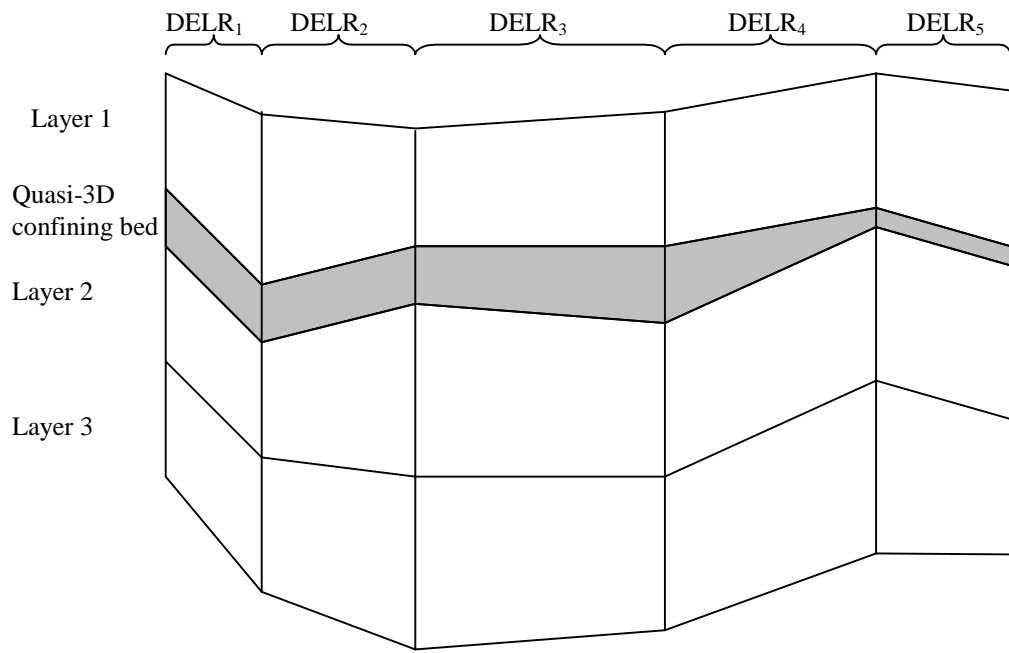
MODFLOW is designed to simulate steady state or transient conditions. For steady state, the storage term in the ground-water flow equation (see the Ground-Water Flow Process Section) is set to zero. This is the only part of the flow equation that depends on length of time, so the stress-period length does not affect the calculated heads in a steady-state simulation. However, it is required in MODFLOW-2000, as in earlier versions of MODFLOW, that the length of a steady-state stress period be specified, partly so that the same input mechanism for all stress periods can be used. A single time step is all that is required for steady-state stress periods. Unlike earlier versions of MODFLOW, the stress period length can be zero, but care is needed because a non-zero length may be important for other processes such as transport (GWT).

The biggest differences in the way stress periods are implemented compared to previous versions of MODFLOW is that MODFLOW-2000 allows individual stress periods in a single simulation to be either transient or steady state instead of requiring the entire simulation to be either steady state or transient. Steady-state and transient stress periods can occur in any order. Commonly the first stress period is steady state and produces a solution that is used as the initial condition for subsequent transient stress periods.

The time-discretization information is included in the new discretization file along with space discretization information. Previously, the time-discretization information was included in the Basic Package file.



(A) Plan View



(B) Cross section along row i .

Figure 2.--Finite-difference grid with (A) plan view and (B) cross-section view.

Units of Length and Time

As in all earlier MODFLOW versions, the Ground-Water Flow Process of MODFLOW-2000 formulates the ground-water flow equation without using prescribed length and time units. Any consistent units of length and time can be used when specifying the input data for a simulation. The user can set flags that specify the length and time units (see the input instructions for the Discretization File), which may be useful in various parts of MODFLOW; however, it is not a requirement to specify which units are being used. For example, the Basic Package uses the time-units flag (ITMUNI) to be able to print a table of simulation time that is labeled with time units. If the time units are not specified, the program still runs, but the table of simulation time does not indicate the time units. A length-unit flag, which was not included in any of the earlier versions of MODFLOW, has been added to MODFLOW-2000. It is expected that other processes will generally work with consistent length and time units; however, there could be situations in which there is a requirement to specify the length or time units. In such situations, the input instructions will state the requirements.

GROUND-WATER FLOW PROCESS

This section describes the Ground-Water Flow Process. The flow equation used in MODFLOW is briefly summarized, followed by a description of how parameters are used. Then the packages that are incorporated are described. In addition to including many previously documented packages, a new package called the Layer-Property Flow (LPF) Package has been added. Complete documentation of LPF is included in this report. Another part of this section describes a program that converts input data for a MODFLOW-96 simulation to data for MODFLOW-2000. The final part describes the ability to add packages to the Ground-Water Flow Process.

Ground-Water Flow Equation

The partial-differential equation of ground-water flow used in MODFLOW is (McDonald and Harbaugh, 1988, p. 2-1)

$$\frac{\partial}{\partial x} \left(K_{xx} \frac{\partial h}{\partial x} \right) + \frac{\partial}{\partial y} \left(K_{yy} \frac{\partial h}{\partial y} \right) + \frac{\partial}{\partial z} \left(K_{zz} \frac{\partial h}{\partial z} \right) + W = S_s \frac{\partial h}{\partial t} \quad (1)$$

where

K_{xx} , K_{yy} , and K_{zz} are values of hydraulic conductivity along the x, y, and z coordinate axes, which are assumed to be parallel to the major axes of hydraulic conductivity (L/T);

h is the potentiometric head (L);

W is a volumetric flux per unit volume representing sources and/or sinks of water, with $W < 0.0$ for flow out of the ground-water system, and $W > 0.0$ for flow in (T^{-1});

S_s is the specific storage of the porous material (L^{-1}); and

t is time (T).

Equation 1, when combined with boundary and initial conditions, describes transient three-dimensional ground-water flow in a heterogeneous and anisotropic medium, provided that the principal axes of hydraulic conductivity are aligned with the coordinate directions.

The Ground-Water Flow Process solves equation 1 using the finite-difference method in which the ground-water flow system is divided into a grid of cells (fig. 2). For each cell, there is a single point, called a node, at which head is calculated. The finite-difference equation for a cell is (McDonald and Harbaugh, 1988, p. 2-18)

$$\begin{aligned}
& CR_{i,j-\frac{1}{2},k} (h_{i,j-1,k}^m - h_{i,j,k}^m) + CR_{i,j+\frac{1}{2},k} (h_{i,j+1,k}^m - h_{i,j,k}^m) \\
& + CC_{i-\frac{1}{2},j,k} (h_{i-1,j,k}^m - h_{i,j,k}^m) + CC_{i+\frac{1}{2},j,k} (h_{i+1,j,k}^m - h_{i,j,k}^m) \\
& + CV_{i,j,k-\frac{1}{2}} (h_{i,j,k-1}^m - h_{i,j,k}^m) + CV_{i,j,k+\frac{1}{2}} (h_{i,j,k+1}^m - h_{i,j,k}^m) \\
& + P_{i,j,k} h_{i,j,k}^m + Q_{i,j,k} = SS_{i,j,k} (\text{DELR}_j \times \text{DEL}C_i \times \text{THICK}_{i,j,k}) \frac{h_{i,j,k}^m - h_{i,j,k}^{m-1}}{t^m - t^{m-1}}
\end{aligned} \tag{2}$$

where

$h_{i,j,k}^m$ is head at cell i,j,k at time step m (L);

CV, CR, and CC are hydraulic conductances, or branch conductances, between node i,j,k and a neighboring node (L^2/T);

$P_{i,j,k}$ is the sum of coefficients of head from source and sink terms (L^2/T);

$Q_{i,j,k}$ is the sum of constants from source and sink terms, with $Q_{i,j,k} < 0.0$ for flow out of the ground-water system, and $Q_{i,j,k} > 0.0$ for flow in (L^3/T);

$SS_{i,j,k}$ is the specific storage (L^{-1});

DELR_j is the cell width of column j in all rows (L);

$\text{DEL}C_i$ is the cell width of row i in all columns (L);

$\text{THICK}_{i,j,k}$ is the vertical thickness of cell i,j,k (L); and

t^m is the time at time step m (T).

To designate hydraulic conductance between nodes, as opposed to hydraulic conductance within a cell, the subscript notation “1/2” is used. For example, $CR_{i,j+1/2,k}$ represents the conductance between nodes i,j,k and $i,j+1,k$.

For steady-state stress periods, the storage term and, therefore, the right-hand side of equation 2, is set to zero.

The application of equation 2 to all cells defines a set of simultaneous equations, and these equations are solved for head at each node. For solution by computer, equation 2 is modified into the form

$$\begin{aligned}
& CV_{i,j,k-\frac{1}{2}} h_{i,j,k-1} + CC_{i-\frac{1}{2},j,k} h_{i-1,j,k} + CR_{i,j-\frac{1}{2},k} h_{i,j-1,k} \\
& + (-CV_{i,j,k-\frac{1}{2}} - CC_{i-\frac{1}{2},j,k} - CR_{i,j-\frac{1}{2},k} - CR_{i,j+\frac{1}{2},k} - CC_{i+\frac{1}{2},j,k} - CV_{i,j,k+\frac{1}{2}} + \text{HCOF}_{i,j,k}) h_{i,j,k} \\
& + CR_{i,j+\frac{1}{2},k} h_{i,j+1,k} + CC_{i+\frac{1}{2},j,k} h_{i+1,j,k} + CV_{i,j,k+\frac{1}{2}} h_{i,j,k+1} = \text{RHS}_{i,j,k} \cdot
\end{aligned} \tag{3}$$

This equation applies to time step m ; however, the time superscript has been removed for simplicity. $\text{HCOF}_{i,j,k}$ contains $P_{i,j,k}$ and the negative of the part of the storage term that includes the head in the current time step m (the negative sign comes from moving the term to the left-hand side). RHS includes $-Q$ (the negative sign comes from moving Q to the right-hand side) and the part of the storage term that is multiplied by the head at time step $m-1$.

The CV, CR, and CC coefficients and the storage-related parts of HCOF and RHS are all calculated by a single package, which is called an internal flow package. Each package that contributes a different source or sink term is called a source-term package. Sinks are viewed as negative sources. The structure of MODFLOW was designed so that any number of source-term packages can be in use in a simulation, but there can be only a single internal-flow package in use. However, there can be multiple internal-flow packages available from which to choose. The original MODFLOW included the Block-Centered Flow (BCF) Package; subsequently, the

Generalized Finite-Difference (GFD) Package was developed (Harbaugh, 1992). This report documents another internal-flow package called the Layer-Property Flow (LPF) Package.

Equation 3 is written only at cells for which head must be calculated. A variable named IBOUND is defined at each cell to indicate that the head in the cell should be calculated (called a variable-head cell); that water cannot flow through the cell (called a no-flow cell); or that the head should not change from a user-specified value (called a constant- or specified-head cell). IBOUND is discussed further in the documentation of the Layer-Property Flow Package.

Use of Parameters in the Ground-Water Flow Process

A new feature of MODFLOW-2000 is that many of the numerous data values that must be specified for each model cell can be specified using parameters. As mentioned earlier, a parameter is a single value that can be used to determine data values for multiple cells. Parameters can often make data input more convenient because of the multi-cell capability. For example, parameters can make it easier to adjust model data when manually calibrating a model or making multiple projection simulations in which many data values must be modified by prescribed amounts. Also, parameters are required when using some other processes. For example, the Parameter-Estimation Process requires parameters to be defined because it is generally impossible to estimate the optimum values for all the types of data at all cells. Even in a small model, there are thousands of data values used by the Ground-Water Flow Process, yet the available observation data are typically only sufficient to estimate a relatively small number of values. Thus, the Parameter-Estimation Process estimates the optimum value for a more limited number of parameters.

The use of parameters in the Ground-Water Flow Process is described in the following sections. The two general data-input structures that must be accommodated through the use of parameters are described first. Then the details of how parameters determine the data value at cells are described. The most common and direct approach for determining data values from parameters is to have the value for an individual cell be defined by one parameter. A more complex approach can also be used in which the data value for a single cell is determined by adding contributions from multiple parameters. This additive approach allows interpolation techniques, such as kriging, and stochastic techniques, such as the pilot-point method, to be used to produce smooth variations of data values throughout a region based upon the multiple parameter values.

Layer Data and List Data

Layer data refers to any type of data for which a value is required for every cell in one or more horizontal layers of the grid. Examples of layer data include areal recharge flux, hydraulic conductivity, and specific storage. In MODFLOW-2000, as in previous versions of MODFLOW, one approach for defining layer data is to directly read it as input data. This direct approach is implemented by utility modules, which provide a common mechanism for reading the layer data required for any package. When layer data are required for multiple layers, the utility models read the data a layer at a time. For each layer, the user can specify either a single value that will apply to all of the cells in the layer or individual values for each cell, which are read row by row starting at row 1. Unlike earlier versions of MODFLOW, however, MODFLOW-2000 allows some layer data to be defined using parameters. The method for defining layer data from parameters is described in the following section. For layer data, the user can usually choose between either directly reading the data (through the utility modules) or using parameters, but the same method must be used consistently for any type of input data. That is, if parameters are used to define hydraulic conductivity in a layer, parameters must be used to define hydraulic conductivity for all layers.

List data refers to any type of data for which data values are required for only some of the cells in the grid. Examples include the well recharge rate as simulated by the Well Package and the riverbed conductance as simulated by the River Package. As with layer data, one approach for defining list data in MODFLOW-2000 is to directly read it as input data. When list data are read by MODFLOW-2000, one line of data is read for each cell for which data are required. For most of the packages that read list data, each line of data includes the layer, row, and column of the cell for which the data applies and one or more types of data. For example, the River Package

requires three types of data for each cell at which a river interacts with the ground-water system (river stage, riverbed conductance and riverbed bottom elevation), and each line of data includes all three data types. MODFLOW-2000 allows some list data to be defined using parameters. The method for defining list data from list parameters is described in the following section. When parameters are allowed for defining list data, it is generally possible to define some values in a list by directly reading and other values in the same list through parameters.

Case 1: One Parameter Is Used to Determine a Cell Data Value

When parameters are used, the data value for a cell is calculated as the product of the parameter value, which might apply to many cells, and a cell multiplier, which applies to only that cell. The multiplier makes it possible for a single parameter to define different data values for different cells associated with the same parameter. For example, consider a riverbed for which the riverbed conductance is to be defined using a parameter. Field data might indicate that the riverbed has a uniform hydraulic conductivity over a length that covers many cells, but the geometry of the riverbed varies from cell to cell. A single parameter can be used to represent the uniform riverbed hydraulic conductivity. The multiplier for each cell associated with the parameter would then represent the area of the riverbed in the cell divided by the thickness of the riverbed. Thus, the final data value for each cell would be the product of the riverbed hydraulic conductivity and riverbed area divided by thickness, which is the riverbed conductance.

Another example is hydraulic conductivity for model cells. Consider a situation in which the field data indicates that in one region of an aquifer, there are interbedded coarse- and fine-grained sediments. The fine-grained sediments have such low hydraulic conductivity that their contribution to the transmissivity is negligible. The coarse-grained sediments are assumed to have a generally uniform hydraulic conductivity, and the proportion of high-conductivity sediments has been mapped throughout the region. To avoid using many parameters to represent the varying average hydraulic conductivity resulting from the combination of coarse and fine materials, a single parameter representing the uniform hydraulic conductivity of the coarse-grained material could be defined. The multipliers for individual cells could then represent the fraction of the total thickness that is coarse-grained material. The product of the parameter and multiplier would then represent the average hydraulic conductivity for each cell.

Parameters for list and layer data require the following information to be specified:

- Type of data to be determined,
- The cells for which data values are to be determined, and
- Multipliers to be used in determining data values.

As described in the following sections, the methods for defining this information is different for layer and list data.

Parameters for List Data

Each package that incorporates list data requires one or more types of data to be defined for listed cells. The input instructions for a package indicate which data types can be defined using parameters. Each data type is given a specific name that must be included as part of the input data that defines a parameter of that type. For example, in the River Package, there are three types of list data (river stage, riverbed conductance, and riverbed bottom elevation), but only riverbed conductance can be defined using parameters. The parameter type for riverbed conductance is RIV.

For list data, the cells that are associated with each parameter are defined in a list in which each line is identical to the line that would normally occur in the package input file with one exception: In the position where the data of the specified type would be, if parameters were not being used, is a multiplier rather than the value to be directly used by the package. The other data values on the line are the same as they would otherwise be, if parameters were not being used.

For example, consider again the use of parameters to define data for the River (RIV) Package. When parameters are not used, input data for a cell includes six values: layer, row, column, stage, riverbed conductance,

and riverbed bottom elevation. When parameters are used, there is a list for each parameter that also contains six values per line. The fifth value is the multiplier for riverbed conductance; the other five values are the same as when parameters are not used. Thus, a non-parameter list can be made into a parameter list by changing just the fifth value on each line. If the parameter value is specified as 1.0, the lines would be identical.

Here is an illustration of how list data are calculated using contributions from two RIV parameters, R1 and R2. The data list for parameter R1 is:

Layer	Row	Column	River Stage	Riverbed Conductance Multiplier	Riverbed Bottom Elevation
1	4	5	4.5	150	4.0
1	5	6	4.8	180	4.3
1	5	7	5.0	200	4.5
1	6	7	5.1	250	4.6

The data list for parameter R2 is:

Layer	Row	Column	River Stage	Riverbed Conductance Multiplier	Riverbed Bottom Elevation
1	10	22	8.9	500	8.0
1	11	21	9.5	600	8.5
1	11	20	10.2	700	9.0

If R1 = 10, and R2 = 20, the resulting list data for the river is:

Layer	Row	Column	River Stage	Riverbed Conductance	Riverbed Bottom Elevation
1	4	5	4.5	1500	4.0
1	5	6	4.8	1800	4.3
1	5	7	5.0	2000	4.5
1	6	7	5.1	2500	4.6
1	10	22	8.9	10000	8.0
1	11	21	9.5	12000	8.5
1	11	20	10.2	14000	9.0

Parameters for Layer Data

Each package that incorporates layer data may have any number of types of data to be defined. The input instructions for a package indicate which data types can be defined using parameters. Each data type is given a specific name that must be included as part of the input data that defines a parameter of that type. For example, in the Layer-Property Flow Package, there are six types of layer data, such as horizontal hydraulic conductivity and specific storage, that can be defined using parameters, and one, called WETDRY, that cannot. The parameter type for horizontal hydraulic conductivity is HK, and the parameter type for specific storage is SS.

For layer data, parameter multipliers are defined using multiplier arrays. “Array” is programming terminology for a variable that contains many values. In this case, each multiplier array contains values for every cell in a layer, and the values can be individually referenced using a row and column index. There can be a different multiplier array for every layer to which the parameter applies, and these are identified when the parameter is defined. Thus, the multiplier for cell (i,j,k) is the (i,j) value of the multiplier array that is specified for layer “k.”

To allow only some of the cells of a layer to be associated with a layer parameter, a capability called zonation is used. Like multiplier arrays, each zone array is named and contains values for every cell in a layer. Values in a zone array are integers. There can be a different zone array for every layer to which the parameter applies. When a parameter is defined, the zone array and one or more zone values are specified. The parameter applies to cells at which the value of the zone array matches any one of the specified zone values; that is, the data value at a cell is the product of the multiplier array at the cell and the parameter value only if the value of the zone array matches one of the zone values specified for the parameter.

Multiplier and zone arrays are defined as part of input to the Global Process. These arrays are read by the array utility modules the same way as layer data for a single layer are read.

As an example of how layer data are calculated from parameters, consider the following multiplier and zone arrays for a layer that has 3 rows and 4 columns.

Multiplier array:

		Column			
		1	2	3	4
Row	1	200.	30.	350.	400.
	2	200.	30.	60.	400.
	3	15.	30.	60.	425.

Zone array:

		Column			
		1	2	3	4
Row	1	1	3	1	1
	2	1	2	3	4
	3	3	3	3	5

If parameter P1 applies to values where the zone array is 2 or 3 and parameter P2 applies to values where the zone array is 1, 4, or 5; then the following input values would be calculated.

		Column			
		1	2	3	4
Row	1	$200 \times P2$	$30 \times P1$	$350 \times P2$	$400 \times P2$
	2	$200 \times P2$	$30 \times P1$	$60 \times P1$	$400 \times P2$
	3	$15 \times P1$	$30 \times P1$	$60 \times P1$	$425 \times P2$

If P1=50 and P2 =80, the final result is:

		Column			
		1	2	3	4
Row	1	16000	1500	28000	32000
	2	16000	1500	3000	32000
	3	750	1500	3000	34000

As already mentioned, a single layer parameter can specify data for multiple layers. For each layer that is included in a parameter, there can be a different multiplier and zone array. The above procedure is applied to calculate the data values for the cells in each layer.

As is shown in the example, the use of a multiplier array allows layer data that is determined by one parameter to be different at each cell. Further, the relative values of the parameter-based layer data are the same as the relative values of the multipliers regardless of the value of the parameter. Consider parameter P1 in the example above. The multiplier for each successive column from left to right is twice the value of the previous column. Thus, the resulting data values that are based on P1 also differ by a factor of 2 from left to right.

Case 2: Additive Parameters Are Used to Determine a Cell Data Value

In the above discussion, it is assumed that the data value for a cell is determined by a single parameter. It is possible, however, for the value at a cell to be determined by more than one parameter. If two or more parameters of the same type include the same cell, the final data value equals the sum of the contributions from all of the parameters, where the contribution from each parameter is calculated as described above. When two or more parameters are used to define the value for cells, the parameters are said to be additive.

As an example of using two parameters to determine the data values for the same cells, assume that it is desired to have the distribution of horizontal hydraulic conductivity, which is layer data, vary linearly from 10 to 100 from the left side to the right side of a layer. This can be done using two additive parameters, P1 and P2, that both apply to all cells in the layer. For a layer consisting of 8 rows and 5 columns, the multiplier array for P1 is:

		Column				
		1	2	3	4	5
Row	1	1.0	0.75	0.50	0.25	0.0
	2	1.0	0.75	0.50	0.25	0.0
	3	1.0	0.75	0.50	0.25	0.0
	4	1.0	0.75	0.50	0.25	0.0
	5	1.0	0.75	0.50	0.25	0.0
	6	1.0	0.75	0.50	0.25	0.0
	7	1.0	0.75	0.50	0.25	0.0
	8	1.0	0.75	0.50	0.25	0.0

The multiplier array for P2 is:

		Column				
		1	2	3	4	5
Row	1	0.0	0.25	0.50	0.75	1.0
	2	0.0	0.25	0.50	0.75	1.0
	3	0.0	0.25	0.50	0.75	1.0
	4	0.0	0.25	0.50	0.75	1.0
	5	0.0	0.25	0.50	0.75	1.0
	6	0.0	0.25	0.50	0.75	1.0
	7	0.0	0.25	0.50	0.75	1.0
	8	0.0	0.25	0.50	0.75	1.0

If P1 = 10., and P2=100., the contributions from both parameters will be added as follows:

The multiplier array for P2 is:

		Column				
		1	2	3	4	5
Row	1	10+0	7.5+25	5+50	2.5+75	0+100
	2	10+0	7.5+25	5+50	2.5+75	0+100
	3	10+0	7.5+25	5+50	2.5+75	0+100
	4	10+0	7.5+25	5+50	2.5+75	0+100
	5	10+0	7.5+25	5+50	2.5+75	0+100
	6	10+0	7.5+25	5+50	2.5+75	0+100
	7	10+0	7.5+25	5+50	2.5+75	0+100
	8	10+0	7.5+25	5+50	2.5+75	0+100

The final result is the desired values:

		Column				
		1	2	3	4	5
Row	1	10.	32.5	55	77.5	100.
	2	10.	32.5	55	77.5	100.
	3	10.	32.5	55	77.5	100.
	4	10.	32.5	55	77.5	100.
	5	10.	32.5	55	77.5	100.
	6	10.	32.5	55	77.5	100.
	7	10.	32.5	55	77.5	100.
	8	10.	32.5	55	77.5	100.

This same distribution could be created in other ways. For example, it could be created without parameters by simply reading the final data values as input data. It could also be generated using a single parameter by having the multiplier vary linearly from 1.0 on the left and 10.0 on the right and having a parameter value be 10.0. There would be an advantage of using two parameters, however, if there were a need to try many distributions that have a linear variation from left to right. The two-parameter approach would allow any such linear distribution to be created by simply changing the two parameter values.

Additive parameters can also be used for list data. Consider another hypothetical situation in which the River Package is being used. In this situation the riverbed sediments are coarse grained at the head of the river and fine grained at the mouth. The resulting riverbed conductance along the first three cells representing the river is assumed to be a constant indicative of the coarse-grained sediments, and the conductance in the last three cells is assumed to be a constant indicative of the fine-grained sediments. In the middle region of the river, which crosses five model cells, it is assumed that the two materials are mixed in such a manner that the riverbed conductance varies linearly from the coarse-grained value to the fine-grained value. This riverbed conductance distribution can be represented by two additive parameters—one for the coarse-grained material and one for the fine-grained material. The following is an example of a data set for this situation in which P1 represents the riverbed conductance of the coarse-grained material and P2 represents the riverbed conductance of the fine-grained material. The data list for parameter P1 is:

Layer	Row	Column	River Stage	Riverbed Conductance Multiplier	Riverbed Bottom Elevation
1	7	9	7.5	1.0	7.0
1	8	9	6.8	1.0	6.3
1	9	9	5.9	1.0	5.4
1	10	9	5.1	.83	4.6
1	11	9	4.5	.67	4.0
1	12	9	3.8	.50	3.3
1	13	9	3.2	.33	2.7
1	14	9	2.5	.17	2.0

The data list for parameter P2 is:

Layer	Row	Column	River Stage	Riverbed Conductance Multiplier	Riverbed Bottom Elevation
1	10	9	5.1	.17	4.6
1	11	9	4.5	.33	4.0
1	12	9	3.8	.50	3.3
1	13	9	3.2	.67	2.7
1	14	9	2.5	.83	2.0
1	15	9	2.1	1.0	1.6
1	16	9	1.8	1.0	1.3
1	17	9	1.5	1.0	1.0

Parameters P1 and P2 both include the five cells where the two riverbed materials are assumed to intermix, so the riverbed conductance for those cells will be the sum of the components from both parameters. For example, the river reach at cell (1, 12, 9) is in the middle of the transition region, and the conductance multipliers are both 0.5 so that the riverbed conductance will be the sum of half of P1 and half of P2.

In addition to the above examples of linear interpolation, additive parameters can be used to generate data using many point-based interpolation techniques, such as distance weighting and kriging. While the regression part of the pilot-point method (RamaRao and others, 1995) could be performed by MODFLOW-2000, the selection of pilot points and redefinition of multiplication arrays for the kriging would need to be done externally.

Packages Included in this Report

This section very briefly describes the current versions of the 14 previously existing packages and one new package that are included in the initial release of MODFLOW-2000 and discussed in this work. As has occurred with previous versions of MODFLOW, other packages and their documentation may be added to U.S. Geological Survey distributions of MODFLOW-2000. The 15 packages discussed in this document include

- the 10 documented in McDonald and Harbaugh (1988),
- the more recent solvers PCG2 (Hill, 1990) and DE4 (Harbaugh, 1995),
- the Hydraulic Flow Barrier (HFB) Package (Hsieh and Freckleton, 1993),
- the Time-Variant Specified Head Package (CHD; Leake and Prudic, 1991), and
- the new Layer Property Flow Package (LPF).

The new Layer-Property Flow Package is completely documented in this report.

Previously Existing Packages

Basic (BAS) Package

The input requirements for the BAS Package data file in MODFLOW-2000 have been modified compared to the MODFLOW-96 version (Harbaugh and McDonald, 1996a) to remove the parts that have been incorporated into the Global Process discretization file. This includes the number of layers, rows, and columns in the grid and the number and length of stress periods. Variables IAPART and ISTRT have also been eliminated from the BAS input file. They were used to conserve memory, but the need to conserve memory on modern computers is not as great as it used to be when MODFLOW was first developed. The elimination of ISTRT has an impact on default output control—drawdown will no longer be printed for default output control, and an output control input file will be needed to obtain drawdowns. All these changes make it necessary to modify an existing MODFLOW-96 BAS file in order to work with MODFLOW-2000. The remaining parts of the input file are the variables HNOFLW, STRT (initial head), and IBOUND. The translation program described below will generate a MODFLOW-2000 BAS file from a MODFLOW-96 BAS file.

The file type for the BAS Package input file in the Name File is BAS6. If a MODFLOW-96 Name File, which would specify a file type of BAS for the BAS input file, is used, MODFLOW-2000 will stop the simulation and print an error message indicating that the file type is illegal. This avoids the input errors that would occur if the MODFLOW-2000 version of BAS were to read a MODFLOW-96 BAS input file.

When the Parameter-Estimation Process is used, the Ground-Water Flow BAS input file is rewound each time the Ground-Water Flow Process is restarted so that the input data will be the same each time. For this reason, the BAS Package writes its output to the LIST file, which is also rewound each time the Ground-Water Flow Process is restarted if there is a GLOBAL file. The LIST file will therefore contain output from the BAS Package for the last execution of the Ground-Water Flow Process.

The Output Control Option of the BAS Package has been modified to allow the boundary flag (variable IBOUND) for every cell to be written into a disk file at the end of any time step. This is useful in simulations in which cells switch between wet and dry, thus changing the value of IBOUND.

Block-Centered Flow (BCF) Package

The BCF Package functions internally exactly as in MODFLOW-96, which means that it includes the original functionality (McDonald and Harbaugh, 1988, ch. 5), the capability to convert dry cells to wet (McDonald and others, 1992), and alternate methods for calculating interblock transmissivity (Goode and Appel, 1992). The MODFLOW-2000 version has been modified, however, so that it no longer reads any discretization data. Instead, it uses the horizontal grid spacing (DELR and DELC), layer elevation (TOP and BOT variables in MODFLOW-96), and transient/steady-state data included in the Global Process discretization file. Therefore, an existing MODFLOW-96 BCF file must be modified to work with MODFLOW-2000. The translation program (MF96TO2K) described below will generate a MODFLOW-2000 BCF file from a MODFLOW-96 BCF file.

The layer elevations defined in the discretization file are not always used by the BCF Package. Specifically, transmissivity and storage coefficient for confined layers, and the vertical leakance (VCONT) for all layers, are directly read by the BCF Package, and these values incorporate layer thickness. Thus, the global elevation data will not have any impact on BCF calculations in which these input data are used. Further, the BCF Package will not make use of the information in the discretization file that defines confining beds that are simulated using the Quasi-3D Approach because this method is handled in BCF by incorporating the effect of the confining bed in the user-specified VCONT values (McDonald and Harbaugh, 1988, p. 5-16). When the BCF Package is active, the global elevation data from the discretization file are used only for the head-dependent formulations: saturated thickness calculation for water-table and convertible layers, storage conversion for convertible layers, and vertical flow limitation that occurs when the hydraulic head in a layer falls beneath the bottom of the layer above.

The parameter method of data input has not been incorporated into the BCF Package. The new Layer-Property Flow Package, which can substitute for BCF, incorporates parameters.

The file type for the BCF Package input file in the name file is BCF6. If a MODFLOW-96 name file, which would specify a file type of BCF for the BCF input file, is used; MODFLOW-2000 will stop the simulation because of an illegal file type. This avoids the input errors that would occur if the MODFLOW-2000 version of BCF were to read a MODFLOW-96 BCF input file. The BCF Package writes all of its output to the LIST file.

Horizontal Flow Barrier (HFB) package

The capability to simulate thin, vertical, and low-permeability geologic features was added to MODFLOW with the introduction of the Horizontal Flow Barrier Package (Hsieh and Freckleton, 1993). As originally written, this package was designed to work with the Block Centered Flow Package. The Horizontal Flow Barrier (HFB) Package has been modified to accommodate the use of parameters and to work with either the Block-Centered Flow Package or the new Layer-Property Flow Package.

The HFB package theory and methodology are unchanged from the original implementation. As described in Hsieh and Freckleton (1993), HFB performs its function by adjusting hydraulic conductance between adjacent cells in a layer. The only changes made for MODFLOW-2000 are in the input requirements and in the support for parameters and parameter estimation. Note that in the terminology of the HFB Package, a barrier is situated on the boundary between two adjacent finite-difference cells in the same layer. Parameters defined in the HFB Package control the hydraulic characteristic of one or more horizontal-flow barriers.

The original version of the HFB Package required the input of the hydraulic characteristic either as barrier transmissivity divided by the width of the horizontal-flow barrier (for layer types 0 and 2 in BCF) or as barrier hydraulic conductivity divided by the width of the horizontal-flow barrier (for layer types 1 and 3 in BCF). In the current HFB Package, the hydraulic characteristic is always the barrier hydraulic conductivity divided by the width of the barrier, regardless of the layer type or flow package (BCF or LPF) used; thus, layer thickness is always used in calculating the contribution to the conductance terms. The HFB Package uses cell elevations specified in the discretization file to calculate cell thickness. Cell thickness is head dependent for layer types 1 and 3 in the BCF Package and for convertible layers in the Layer-Property Flow Package.

The file type for the HFB Package input file in the Name File is HFB6. If a Name File lists a file of type HFB, MODFLOW-2000 will stop the simulation with an error message indicating that an illegal file type was specified. If this error message is encountered, the user should ensure that the file associated with this file type is compatible with the current version of the HFB Package and change the file type to HFB6. Input errors would occur if the MODFLOW-2000 version of HFB were to read an input file prepared for the original version of HFB.

Source-Term Packages

These packages are divided according to whether they are head dependent or head independent. They have all been modified to incorporate parameters. Other than the addition of parameters, these packages function the same as in MODFLOW-96. Input files for MODFLOW-96 will still work in MODFLOW-2000. All of the output is written to the LIST file.

Head-Dependent Packages

- RIV—The riverbed conductance can be defined using parameters.
- DRN—The drain conductance can be defined using parameters.
- GHB—The boundary conductance can be defined using parameters.
- EVT—The maximum evapotranspiration flux can be defined using parameters.

McDonald and Harbaugh (1988) document the concepts of the RIV, DRN, GHB, and EVT Packages.

Head-Independent Packages

- WEL—The well recharge rate (negative sign indicates discharge) can be defined using parameters.
- RCH—The Recharge flux can be defined using parameters.

McDonald and Harbaugh (1988) document the concepts of both of these packages.

Time-Variant Specified-Head Package

MODFLOW-2000 contains the Time-Variant Specified-Head (CHD) Package (Leake and Prudic, 1991), and this has been modified to allow parameters to define the specified head. Input files for MODFLOW-96 will still work in MODFLOW-2000. All of the output is written to the LIST file.

Solver Packages

Four solvers are included in MODFLOW-2000:

- SIP—strongly implicit procedure (McDonald and Harbaugh, 1988),
- SOR—slice successive overrelaxation (McDonald and Harbaugh, 1988),
- PCG—preconditioned Conjugate Gradient (Hill, 1990), and
- DE4—direct solution based on alternating diagonal ordering (Harbaugh, 1995).

There is no fundamental change to the solvers in MODFLOW-2000. From the perspective of the solvers, the information that defines the simultaneous equations that must be solved at each cell is exactly the same as in earlier versions of MODFLOW. The solvers were modified superficially to allow them to be used by the Sensitivity Process. The user-specified input data to control the solvers is unchanged from that used in MODFLOW-96, except for a minor change for the PCG Package. A new option has been added to the PCG Package, which outputs solver convergence information only if the solver fails to meet closure criteria. Otherwise, no convergence information is written. Input files for MODFLOW-96 will still work in MODFLOW-2000. Package identification and information that confirms what was read from the solver package input file is written to the GLOBAL file. Convergence information is written to the LIST file.

New Package: Layer-Property Flow Package

Introduction

The Layer-Property Flow (LPF) Package is an internal flow package, so, as previously mentioned, it calculates the CV, CR, and CC conductance coefficients and the parts of HCOF and RHS (equation 3) dealing with ground-water storage. The BCF Package also performs this same function, and accordingly, LPF should be viewed as an alternative to the BCF Package. That is, both packages should not be used simultaneously. The following documentation of the LPF Package highlights the similarities and differences between the two packages.

When calculating conductance coefficients, it is assumed in the LPF Package that a node is located at the center of each model cell. The same assumption is also used in the BCF Package, and accordingly, the BCF and LPF Packages are conceptually quite similar. The differences are primarily in the input data that the user specifies. All the input data that define hydraulic properties are independent of cell dimensions in LPF, whereas some of the input data for the BCF Package incorporate cell dimensions. For example, in some situations, the user specifies transmissivity as input data for the BCF Package, but LPF never reads transmissivity. Instead, LPF always reads hydraulic conductivity (either directly or using parameters) and calculates transmissivity by using cell thickness that is determined from the global elevation data. Similarly, vertical leakance (hydraulic conductivity divided by distance between nodes) is directly read by the BCF Package whereas the LPF Package calculates leakance from vertical hydraulic conductivity and distance between nodes, which is calculated from the global elevation data.

Basic Hydraulic Conductance Equations

The concept of hydraulic conductance (shortened to conductance to save space in this report) is defined in McDonald and Harbaugh (1988, chs. 2 and 5). It is reviewed here, including the calculation of equivalent conductance for elements arranged in series. Conductance is a combination of several parameters used in Darcy's law. Darcy's law defines one-dimensional flow in a prism of porous material (fig. 3) as

$$Q = \frac{-KA(h_2 - h_1)}{L} \quad (4)$$

where

Q is the volumetric flow (L^3T^{-1});

K is the hydraulic conductivity of the material in the direction of flow (LT^{-1});

A is the cross-sectional area perpendicular to the flow (L^2);

h_1-h_2 is the head difference across the prism parallel to flow (L); and

L is the length of the prism parallel to the flow path (L).

Conductance, C, is defined as

$$C = \frac{KA}{L} \quad (5)$$

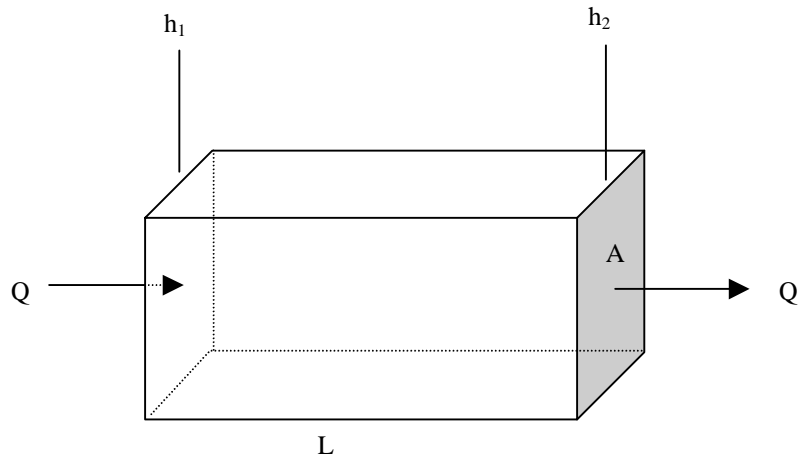


Figure 3.--Prism of porous material illustrating Darcy's law.

Therefore, Darcy's law can be written as

$$Q = C(h_1 - h_2) \quad (6)$$

Another form of the conductance definition for horizontal flow in a prism is

$$C = \frac{TW}{L} \quad (7)$$

where

T is transmissivity (K times thickness of the prism) in the direction of flow (L^2T^{-1}); and

W is the width of the prism (L).

Conductance is defined for a particular prism of material and for a particular direction of flow. In an anisotropic medium that is characterized by three principal directions of hydraulic conductivity, the conductances of a prism in these three principal directions will generally differ.

If a prism of porous material consists of two or more subprisms in series (aligned sequentially in the direction of flow) as shown in figure 4, and the conductance of each subprism is known, a conductance representing the entire prism can be calculated. The equivalent conductance for the entire prism is the rate of flow in the prism divided by the head change across the prism.

$$C = \frac{Q}{h_A - h_B} \quad (8)$$

Assuming continuity of head across each section gives the identity

$$\sum_{i=1}^n \Delta h_i = h_A - h_B \quad (9)$$

Substituting for head change across each section using Darcy's law (equation 6) gives

$$\sum_{i=1}^n \frac{q_i}{C_i} = h_A - h_B \quad (10)$$

Because flow is one dimensional, and we are assuming no accumulation or depletion in storage, all q_i are equal to the total flow Q ; therefore,

$$Q \sum_{i=1}^n \frac{1}{C_i} = h_A - h_B \quad \text{and} \quad \frac{h_A - h_B}{Q} = \sum_{i=1}^n \frac{1}{C_i} \quad (11)$$

By comparing equation 11 with equation 8, it can be seen that

$$\frac{1}{C} = \sum_{i=1}^n \frac{1}{C_i} \quad (12)$$

Thus for a set of conductances arranged in series, the inverse of the equivalent conductance equals the sum of the inverses of the individual conductances. When there are only two sections, the equivalent conductance reduces to

$$C = \frac{C_1 C_2}{C_1 + C_2} \quad (13)$$

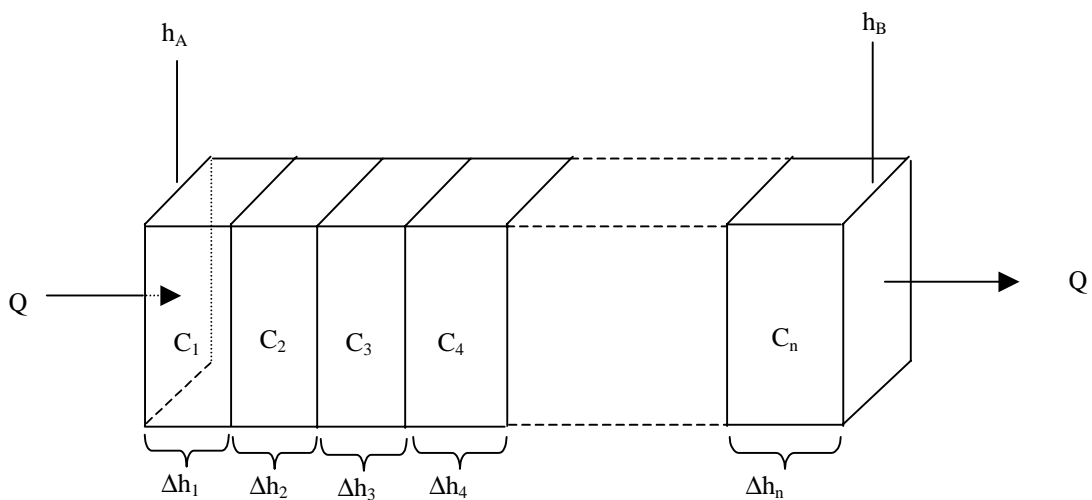


Figure 4.-- Calculation of conductance through several prisms in series.

Horizontal Conductance

The finite-difference equations in MODFLOW use equivalent conductances between nodes of adjacent cells—that is, “branch conductances”—rather than conductances defined within individual cells. The horizontal conductance terms, CR and CC of equation 2, are calculated between adjacent horizontal nodes. CR terms are oriented along rows and thus specify conductance between two nodes in the same row. Similarly, CC terms specify conductance between two nodes in the same column. (Remember that the “1/2” subscript denotes conductance between nodes, as opposed to conductance within a cell. For example, $CR_{i,j+1/2,k}$ represents the conductance between nodes i,j,k and $i,j+1,k$.)

The LPF Package reads data defining the horizontal hydraulic conductivity for individual cells and calculates conductance between nodes. Three methods of calculating these conductances are supported. The methods differ in the assumptions about the way the ground-water system varies from cell to cell.

Constant Product of Hydraulic Conductivity and Thickness within a Cell

The first method is based on the assumption that hydraulic conductivity times thickness, that is, transmissivity, is constant within a cell. There can, therefore, be a discrete change in transmissivity at the boundary between any two cells. By use of this assumption and the previously mentioned assumption in LPF that nodes are in the center of cells, the conductance between the nodes is the equivalent conductance of two half cells in series. Figure 5 illustrates two cells along a row based on these assumptions. Substituting the conductance for each half cell (equation 7) into equation 13 gives:

$$CR_{i,j+1/2,k} = \frac{\frac{TR_{i,j,k} DELC_i}{(1/2)DELR_j} \quad \frac{TR_{i,j+1,k} DELC_i}{(1/2)DELR_{j+1}}}{\frac{TR_{i,j,k} DELC_i}{(1/2)DELR_j} + \frac{TR_{i,j+1,k} DELC_i}{(1/2)DELR_{j+1}}} \quad (14)$$

where

- $TR_{i,j,k}$ is transmissivity in the row direction at cell i,j,k (L^2t^{-1});
- $DELR_j$ is the grid width of column j (L); and
- $DELC_i$ is the grid width of row i (L).

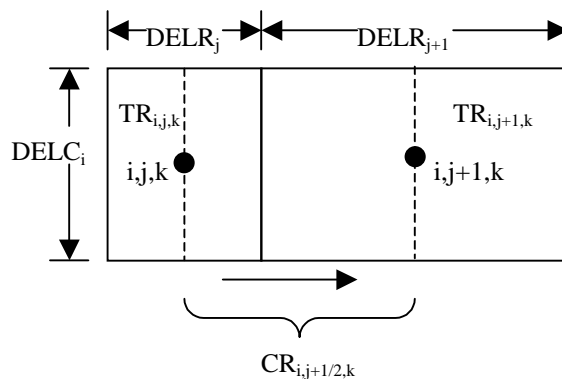


Figure 5.-- Calculation of conductance between nodes using transmissivities and dimensions of cells.

Simplification of equation 14 gives the final equation:

$$CR_{i,j+\frac{1}{2},k} = 2 \text{ DELC}_i \frac{TR_{i,j,k} TR_{i,j+1,k}}{TR_{i,j,k} \text{ DELR}_{j+1} + TR_{i,j+1,k} \text{ DELR}_j} \quad (15)$$

The same process can be applied to the calculation of $CC_{i+1/2,j,k}$ giving:

$$CC_{i+\frac{1}{2},j,k} = 2 \text{ DELR}_i \frac{TC_{i,j,k} TC_{i+1,j,k}}{TC_{i,j,k} \text{ DELC}_{i+1} + TC_{i+1,j,k} \text{ DELC}_i} \quad (16)$$

where

$TC_{i,j,k}$ is the transmissivity in the column direction at cell i,j,k (L^2T^{-1}).

Equations 15 and 16 are used unless the transmissivity of both cells is zero, in which case the conductance between the nodes is set equal to zero without invoking the equations.

Transmissivity is calculated as:

$$TR_{i,j,k} = \text{THICK}_{i,j,k} \text{ HK}_{i,j,k} \quad (17A)$$

$$TC_{i,j,k} = \text{THICK}_{i,j,k} \text{ HK}_{i,j,k} \text{ HANI}_{i,j,k} \quad (17B)$$

where

$\text{HK}_{i,j,k}$ is the hydraulic conductivity of cell i,j,k in the row direction (LT^{-1}),

$\text{HANI}_{i,j,k}$ is the ratio of hydraulic conductivity along columns to the hydraulic conductivity along rows (dimensionless), and

$\text{THICK}_{i,j,k}$ is the saturated thickness of cell i,j,k (L).

$\text{HK}_{i,j,k}$ and $\text{HANI}_{i,j,k}$ are specified as input data. Values of $\text{THICK}_{i,j,k}$ are calculated using cell elevations in the discretization file. The calculation of thickness depends on whether the layer is designated as confined or convertible. This designation is made through the layer type-flag, **LAYTYP**, which is discussed in the Data Requirements Section. In a model layer that is designated as confined, the thickness is the cell thickness:

$$\text{THICK}_{i,j,k} = (\text{TOP}_{i,j,k} - \text{BOT}_{i,j,k}) \quad (18)$$

where

$\text{TOP}_{i,j,k}$ is the elevation of the top of cell i,j,k (L); and

$\text{BOT}_{i,j,k}$ is the elevation of the bottom of cell i,j,k (L).

TOP and BOT are determined from data in the discretization file.

If a layer is designated as convertible, then saturated thickness is calculated throughout the simulation based upon the head ($HNEW_{i,j,k}$):

$$\text{if } HNEW_{i,j,k} \geq TOP_{i,j,k}, \quad \text{then } THICK_{i,j,k} = (TOP_{i,j,k} - BOT_{i,j,k}); \quad (19A)$$

$$\text{if } TOP_{i,j,k} > HNEW_{i,j,k} > BOT_{i,j,k}, \quad \text{then } THICK_{i,j,k} = (HNEW_{i,j,k} - BOT_{i,j,k}); \quad (19B)$$

$$\text{if } HNEW_{i,j,k} \leq BOT_{i,j,k}, \quad \text{then } THICK_{i,j,k} = 0. \quad (19C)$$

At the start of every iteration for solving the flow equation, cell transmissivity values (TR and TC) are recalculated as the product of hydraulic conductivity and saturated thickness (equation 14); then conductance is recalculated using equations 15 and 16. If head drops below the aquifer bottom (equation 19C), the cell is considered to be fully dewatered, and is set to no flow by changing $IBOUND_{i,j,k}$ to 0. The model also has provision for the resaturation of fully dewatered cells, which is described in another section of this report.

Note that unlike the LPF Package, the BCF Package has a special layer type for a water table in layer 1. This BCF layer type always uses equation 19B for calculating saturated thickness. In LPF, the convertible formulation should be used if layer 1 represents a water table. Also, the elevation of the top of layer 1, which is specified in the discretization file, should be higher than the water level will reach at any time in the simulation. This will result in the use of equation 19B for calculating saturated thickness. It is the user's responsibility in this situation to check to see that the simulated water level has not exceeded the top elevation, which would trigger the use of equation 19C for calculating saturated thickness. A logical approach for specifying the top elevation for a layer-1 water table is to set it equal to land-surface elevation. That is, in most situations, it is not expected that the water table will exceed land-surface elevation.

The above approach for calculating interblock conductance is called the "harmonic mean" method. The reason for this name can be seen by manipulating equation 15 into the following:

$$CR_{i,j+\frac{1}{2},k} = \left(\frac{\frac{(1/2)DELR_j + (1/2)DELR_{j+1}}{T_{i,j,k}} + \frac{(1/2)DELR_j + (1/2)DELR_{j+1}}{T_{i,j+1,k}}}{(1/2)DELR_j + (1/2)DELR_{j+1}} \right) \frac{DELC_i}{(1/2)DELR_j + (1/2)DELR_{j+1}} \quad (20)$$

The term in parentheses is the distance weighted harmonic mean of transmissivity of the two half cells, which by comparison with equation 7, can be seen to be the equivalent transmissivity between nodes i,j,k and $i,j+1,k$.

Alternative Approaches for Calculating Horizontal Branch Conductances

Goode and Appel (1992) discuss three alternative approaches for calculating the horizontal branch conductances. These alternative approaches are each based on different assumptions about the flow system: (1) transmissivity varies linearly between nodes, (2) flat, homogeneous aquifer with a water table, (3) flat aquifer with a water table in which hydraulic conductivity varies linearly between nodes. For each approach, a formulation of interblock transmissivity is developed. The interblock conductance is then computed as the interblock transmissivity times cell width divided by distance between nodes according to equation 7.

When transmissivity varies linearly between nodes, the interblock transmissivity derived by Goode and Appel (1992) is:

$$\frac{T_2 - T_1}{\ln(T_2/T_1)} \quad (21)$$

where subscripts 2 and 1 indicate any two adjacent cells along a row or column. This is called the “logarithmic mean” interblock transmissivity method. This formulation will produce exact solutions for steady-state one-dimensional flow if flow is uniform and aligned with the direction of changing transmissivity. To save numerical effort, this function is approximated by

$$\frac{T_1 + T_2}{2} \text{ when } 0.995 \leq \frac{T_2}{T_1} \leq 1.005 .$$

For an unconfined homogeneous aquifer with a flat bottom, the interblock transmissivity derived by Goode and Appel (1992) is:

$$\frac{T_1 + T_2}{2} . \tag{22}$$

This is called the “arithmetic mean” interblock transmissivity method. This formulation will produce exact solutions for uniform, steady-state, one-dimensional flow.

For an unconfined aquifer with a flat bottom and with hydraulic conductivity varying linearly between nodes, Goode and Appel derive the interblock transmissivity as:

$$\left(\frac{\text{THICK}_1 + \text{THICK}_2}{2} \right) \left(\frac{K_1 - K_2}{\ln(K_2/K_1)} \right) . \tag{23}$$

This is called the “arithmetic mean thickness and logarithmic-mean hydraulic conductivity” method. This formulation produces exact solutions for uniform, steady-state, one-dimensional flow when the hydraulic conductivity varies in the direction of flow. To save numerical effort, the hydraulic conductivity part of this function is approximated by $\frac{K_1 + K_2}{2}$ when $0.995 \leq \frac{K_2}{K_1} \leq 1.005$.

Because equation 23 reduces to equation 22 when the aquifer is homogeneous, equation 22 is not separately included in the LPF Package. That is, only equations 21 and 23 are included in the LPF Package along with the harmonic mean method.

Vertical Conductance

Vertical conductance is calculated assuming that nodes are in the center of cells and that there can be discrete changes in vertical hydraulic conductivity at layer boundaries. Figure 6 illustrates this situation for two cells in layers k and k+1. Under these assumptions, the vertical conductance between two nodes will be the equivalent conductance of two half cells in series. Substituting the conductance for each half cell from equation 5 into equation 12 gives

$$\frac{1}{\text{CV}_{i,j,k+\frac{1}{2}}} = \frac{1}{\frac{\text{DELR}_j \text{ DELC}_i \text{ VK}_{i,j,k}}{(\frac{1}{2})\text{THICK}_{i,j,k}}} + \frac{1}{\frac{\text{DELR}_j \text{ DELC}_i \text{ VK}_{i,j,k+1}}{(\frac{1}{2})\text{THICK}_{i,j,k+1}}} \tag{24}$$

where

$\text{VK}_{i,j,k}$ is vertical hydraulic conductivity of cell i,j,k and
 $\text{THICK}_{i,j,k}$ is the saturated thickness of cell i,j,k as defined by equation 18 for confined cells and equation 16 for convertible cells.

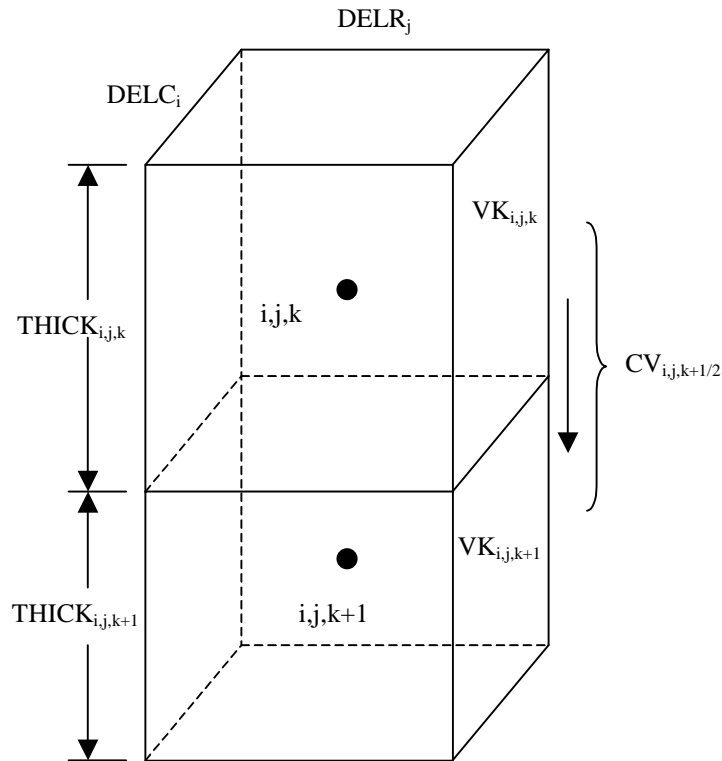


Figure 6.-- Calculation of vertical conductance between two nodes.

Simplification of equation 24 gives

$$CV_{i,j,k+1/2} = \frac{DELR_j DELC_i}{\frac{(1/2)THICK_{i,j,k}}{VK_{i,j,k}} + \frac{(1/2)THICK_{i,j,k+1}}{VK_{i,j,k+1}}} \quad (25)$$

Figure 7 shows a second situation, in which cells i,j,k and $i,j,k+1$ are separated by a semiconfining unit. If the assumptions are made that the semiconfining unit makes no measurable contribution to the horizontal conductance or the storage capacity of either model layer, then the only effect of the confining bed is to restrict vertical flow between the model cells. Under these assumptions, the impact of the semiconfining unit can be simulated without using a separate layer in the finite-difference grid. This is done by including the semiconfining unit in the calculation of vertical conductance between the nodes. As mentioned previously, this is commonly referred to as the "Quasi-Three-Dimensional" (QUASI-3D) Approach (McDonald and Harbaugh, 1988, p. 5-18).

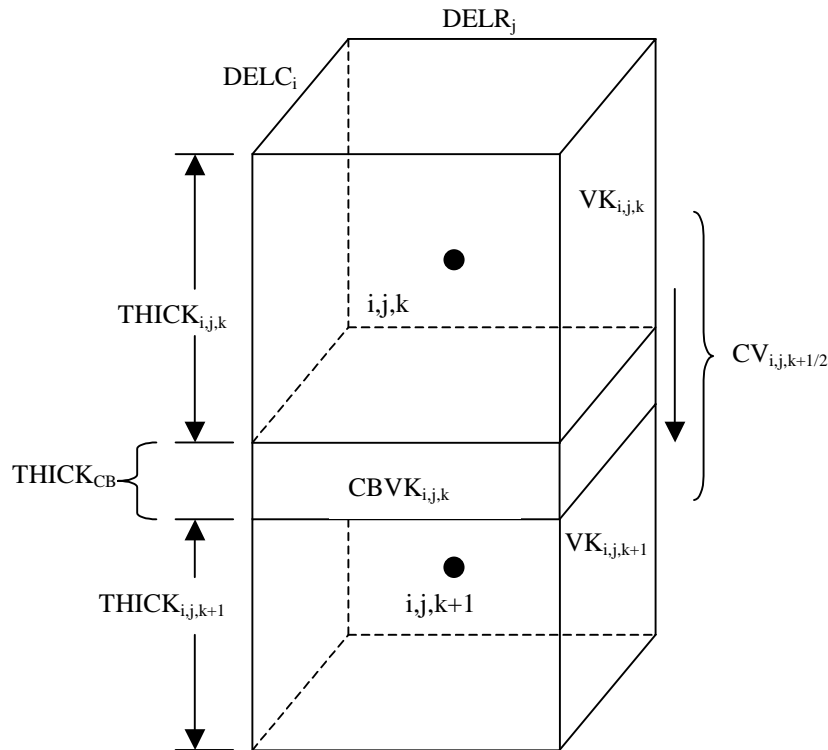


Figure 7.-- Calculation of vertical conductance between two nodes with a semiconfining unit between.

In the Quasi-3D Approach, three intervals must be represented in the summation of conductance between the nodes—the lower half of the upper aquifer, the semiconfining unit, and the upper half of the lower aquifer. Equation 12 can be used to write an expression for the inverse of vertical conductance:

$$\frac{1}{CV_{i,j,k+\frac{1}{2}}} = \frac{1}{\frac{DEL R_j DELC_i VK_{i,j,k}}{(\frac{1}{2})THICK_{i,j,k}}} + \frac{1}{\frac{DEL R_j DELC_i VK_{CB}}{THICK_{CB}}} + \frac{1}{\frac{DEL R_j DELC_i VK_{i,j,k+1}}{(\frac{1}{2})THICK_{i,j,k+1}}} \quad (26)$$

where

VK_{CB} is the hydraulic conductivity of the semiconfining unit between cell i,j,k and $i,j,k+1$, and $THICK_{CB}$ is the thickness of the semiconfining unit.

$THICK_{CB}$ is determined from information in the discretization file, and VK_{CB} is input data for the LPF Package.

Simplification of equation 26 produces

$$CV = \frac{DEL R \ DEL C}{\frac{(1/2)THICK_{i,j,k}}{VK_{i,j,k}} + \frac{THICK_{CB}}{VK_{CB}} + \frac{(1/2)THICK_{i,j,k+1}}{VK_{i,j,k+1}}} \quad (27)$$

Vertical Flow Calculation Under Dewatered Conditions

This section describes a modification to the vertical flow calculation that is applied when a cell in a convertible layer is unconfined (head is below its top elevation), while the cell immediately above is fully or partially saturated. (Convertible layers are specified through the layer type-flag, LAYTYP, which is described in the Data Requirements Section.) The vertical flow correction is applied for every solver iteration.

From equation 2, the term that represents the flow into cell i,j,k through its lower face, which will be called $q_{i,j,k+1/2}$, is

$$q_{i,j,k+1/2} = CV_{i,j,k+1/2} (h_{i,j,k+1} - h_{i,j,k}) \quad (28)$$

where the m superscript that indicates the time step has been removed. Following the convention of MODFLOW, a positive value of $q_{i,j,k+1/2}$ indicates flow into cell i,j,k , and a negative value indicates flow out of the cell. Equations 2 and 28 are based on the assumption that cells i,j,k and $i,j,k+1$ are fully saturated—that is, that the hydraulic head in each cell stands higher than the elevation of the top of the cell.

There are, however, situations in which a part of a confined aquifer may become unsaturated—for example, when drawdown due to pumpage causes water levels to fall, at least locally, below the top of an aquifer. This condition is most likely to occur when an aquifer is overlain by a lower conductivity unit. In terms of simulation, this condition is shown in figure 8. Cells in two model layers are represented. The upper cell (i,j,k) has lower hydraulic conductivity than the underlying cell ($i,j,k+1$), which represents an aquifer. Pumping from the lower layer has lowered the water level in cell $i,j,k+1$ below the elevation of the top of the cell; however, cell i,j,k remains fully saturated. Thus, cell $i,j,k+1$ is effectively unconfined even though the cell above is saturated.

Consider the calculation of flow between nodes in the upper and lower cells. In the upper cell, head is simply whatever head is at that cell— $h_{i,j,k}$. Just below the upper cell, however, unsaturated conditions prevail, so that the pressure sensed on the lower surface of the confining unit is atmospheric—taken as zero in the model formulation. Thus the head at the bottom of the upper cell is simply the elevation at that point—that is, the elevation of the top of the lower cell. If this elevation is designated $TOP_{i,j,k+1}$, then the actual flow through the confining bed is obtained by substituting $TOP_{i,j,k+1}$ for $h_{i,j,k+1}$ in equation 28,

$$q_{i,j,k+1/2} = CV_{i,j,k+1/2} (TOP_{i,j,k+1} - h_{i,j,k}) \quad (29)$$

Thus, the flow will be downward, from cell i,j,k to cell $i,j,k+1$, but under this condition the flow will no longer be dependent on the water level, $h_{i,j,k+1}$, in the lower cell. The simplest approach to this problem in formulating the equation for cell i,j,k would be to substitute the flow expression of equation 29 into equation 2, in place of the expression given in equation 28. However, if we consider the matrix of coefficients of the entire system of finite-difference equations, this direct substitution would render this matrix asymmetric, generating problems in the solution process.

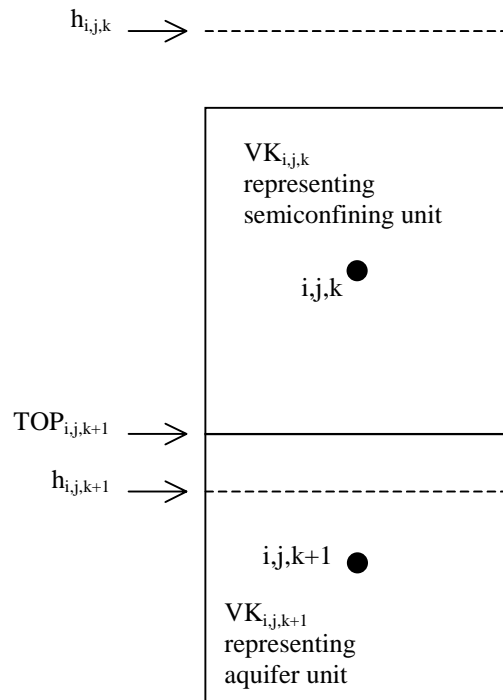


Figure 8.-- Situation in which a correction is required to limit the downward flow into cell $i,j,k+1$, as a result of partial desaturation of the cell.

To avoid this condition, an alternative approach is used. The flow term of equation 28 is allowed to remain on the left side of equation 2. The "actual" flow into cell i,j,k is given by equation 29, so a correction term, q_c , can be obtained by subtracting equation 29 from equation 28:

$$\begin{aligned}
 q_c &= (\text{computed flow into cell } i,j,k) - (\text{"actual" flow into cell } i,j,k) \\
 &= CV_{i,j,k+1/2}(h_{i,j,k+1} - TOP_{i,j,k+1}) .
 \end{aligned}
 \tag{30}$$

This correction term, q_c , should be added to the right side of equation 2 to compensate for allowing the computed flow to remain on the left side of equation 2. This again introduces a problem of matrix unsymmetry because q_c contains the term $h_{i,j,k+1}$. To circumvent this difficulty, q_c is actually computed using the value of $h_{i,j,k+1}$ from the preceding iteration, rather than that from the current iteration:

$$q_c^n = CV_{i,j,k+1/2} (h_{i,j,k+1}^{n-1} - TOP_{i,j,k+1})
 \tag{31}$$

where

q_c^n is the value of q_c to be added to the right side of equation 2 in the n th iteration, and

$h_{i,j,k+1}^{n-1}$ is the value of $h_{i,j,k+1}$ from the preceding iteration, $n-1$.

As convergence is approached, the difference between $h_{i,j,k+1}^{n-1}$ and $h_{i,j,k+1}^n$ becomes progressively smaller, and the approximation involved in equation 31 thus becomes more accurate. In the first iteration of each time step, the initial trial value of $h_{i,j,k+1}$ is used in computing q_c . In terms of equation 3, which is the form of the flow equation that is directly implemented in MODFLOW, q_c^n is added to $RHS_{i,j,k}$.

The process described above is used in formulating the equations for cell i,j,k when the underlying cell, $i,j,k+1$, has "dewatered"—that is, when the water level in $i,j,k+1$ has declined below the top of the cell. A correction must also be applied in formulating the equations for the dewatered cell itself. To compute this correction, cell i,j,k is considered to be the dewatered cell, and we consider flow into i,j,k from the overlying cell, $i,j,k-1$. For this case, the computed flow into cell i,j,k from above is

$$CV_{i,j,k-1/2}(h_{i,j,k-1} - h_{i,j,k}) .$$

whereas the "actual" flow into the cell is

$$CV_{i,j,k-1/2}(h_{i,j,k-1} - TOP_{i,j,k}) .$$

The difference, computed minus "actual" flow, is thus

$$q_c' = CV_{i,j,k-1/2} (TOP_{i,j,k} - h_{i,j,k}) \quad (32)$$

where q_c' is added to the right-hand side of equation 2. From a programming point of view, the most efficient way to handle this correction is to add the term $CV_{i,j,k-1/2}$ to HCOF on the left side of equation 2, while adding the term $CV_{i,j,k-1/2} TOP_{i,j,k}$ to the RHS term. However, this causes difficulties for some solution methods because it reduces the dominance of the diagonal of the coefficient matrix of the finite-difference equations. So the same approach is used as described previously for the vertical correction for the cell above the dewatered cell. That is, q_c' is computed on the basis of the head from the previous iteration, and this correction is added to RHS of equation 3.

This conceptualization of limiting vertical flow is also valid when the unsaturated region in a lower layer is caused by an overlying confining unit that is simulated using the Quasi-3D Approach. Although the confining unit is not simulated as a separate model layer when using this approach, the impact of the confining unit is still simulated through the vertical conductance calculation. The above vertical flow calculation correction is applied. The value of $TOP_{i,j,k}$ is the bottom of the confining bed.

Note that when vertical leakage is limited, an argument could be made that the vertical conductance calculation should also be modified. That is, the vertical conductance should not include any contribution from the lower cell that is partially dewatered. However, this modification of vertical conductance has not been incorporated into this version of the program.

In summary, whenever dewatering of a cell occurs below a cell that is not fully dewatered, two corrections must be made—one in formulating equation 2 as it applies to the overlying cell, and one in formulating equation 2 as it applies to the dewatered cell itself. These two corrections are discussed separately above, in each case using the designation i,j,k to represent the cell for which equation 2 is formulated. It is important to keep in mind, however, that both corrections are applied in any dewatering event, and that the form of the corrections has been

developed to preserve the symmetry of the coefficient matrix of the finite-difference equations and its diagonal dominance. This correction is applied as described for layers that are designated as convertible layers.

Conversion from Dry to Wet

When saturated thickness is 0 as defined by head being less than the bottom elevation of a cell (equation 19C), it is clear that a variable-head (wet) cell should convert to dry. Unfortunately, there is not a straight-forward way to know when a dry cell should convert to wet. McDonald and others (1992) describe an approach for doing this and its application in the BCF Package of MODFLOW. The same approach is applied to the LPF Package. The following is a brief overview of the approach. Readers should refer to McDonald and others (1992) for additional details. In particular, that report describes detailed information about numerical instabilities that can result from using this option, and several test problems are included.

A dry cell converts to wet based upon the head in an adjacent cell compared to a wetting threshold, THRESH, for the cell. If the head at an adjacent cell equals or exceeds the threshold at the start of a solution iteration, the dry cell is converted to wet. The only cells that can cause a dry cell to convert to wet are the four cells that are immediately adjacent to the cell horizontally and the cell immediately below. There is also an option to use only the cell below to determine that a dry cell becomes wet. The head below can be a better wetting indicator than the head at horizontally adjacent cells when the head variations between adjacent horizontal cells are larger than the vertical head variations, which is frequently the case.

A primary difficulty with this approach is that the value of THRESH must be determined by trial and error. If THRESH is too low, a cell may be incorrectly converted to wet. That is, a cell may convert to wet and then revert to dry in later solver iterations because the head does not stay above the bottom elevation. This cycle of converting between wet and dry can repeat indefinitely, preventing the solver from converging. Larger values of THRESH can avoid repeated conversion between wet and dry, but they increase the nonuniqueness of the solution (McDonald and others, 1992, p. 8).

When a cell converts to wet, the initial estimate of head is established according to one of two equations:

$$h_{i,j,k} = BOT_{i,j,k} + WETFCT(h_n - BOT_{i,j,k}) \quad (33A)$$

$$h_{i,j,k} = BOT_{i,j,k} + WETFCT(THRESH_{i,j,k}) \quad (33B)$$

where

WETFCT is a user-specified constant, generally between 0 and 1, and

h_n is the head at the neighboring cell that causes cell i,j,k to convert to wet.

The user chooses the equation that works best for the problem being simulated. The initial estimate of head at a cell that converts to wet is theoretically unimportant because the solver should calculate the correct value in subsequent iterations. In practice, however, the initial estimate is often important for solver efficiency; that is, if a poor choice is made, the solver convergence may be slower than optimum. A bad choice for initial head can also cause the cell to oscillate between wet and dry.

Another user-specified option is the solver iteration interval for attempting to wet cells. For example, if the interval is 4, cell wetting is attempted every 4 iterations. This can prevent the normal fluctuations in heads that can occur in iterative solvers in the next few iterations after cells are converted to wet from inappropriately triggering more cell wetting. All of these options are included to help make the solution process more stable. Refer to McDonald and others (1992) for more information about solution instability and how to deal with it.

Storage Formulation

The LPF Package formulates storage terms for transient stress periods much as done for the BCF Package (McDonald and Harbaugh, 1988, p. 5-24). LPF makes the same distinction between confined layers for which storage terms remain constant throughout the simulation, and those in which the storage terms may "convert"

from a confined value to a water table value, or conversely, as the water level in a cell falls below or rises above the top of the cell. In LPF, this distinction is made through the layer-type flag, LAYTYP, as described in the Data Requirements Section.

For a confined layer, the storage formulation is based upon a direct application of the storage expression in equation 2. This expression for the rate of accumulation of water into a single cell is:

$$\frac{\Delta V}{\Delta t} = SS_{i,j,k} (\text{DELR}_j \text{ DELC}_i \text{ THICK}_{i,j,k}) \frac{h_{i,j,k}^m - h_{i,j,k}^{m-1}}{t^m - t^{m-1}} \quad (34)$$

where

- $SS_{i,j,k}$ is the specific storage of the material in cell i,j,k (L^{-1});
- $\text{THICK}_{i,j,k}$ is the cell thickness as defined by equation 18 (L);
- $h_{i,j,k}^m$ is the head in cell i,j,k at the end of time step m (L);
- $h_{i,j,k}^{m-1}$ is the head in cell i,j,k at the end of time step $m-1$ (L);
- t^m is the time at the end of time step m (T); and
- t^{m-1} is the time at the end of time step $m-1$ (T).

The notation $SC1_{i,j,k}$ is introduced where $SC1_{i,j,k} = SS_{i,j,k} (\text{DELR}_j \text{ DELC}_i \text{ THICK}_{i,j,k})$. In this report the term $SC1_{i,j,k}$ is termed the "storage capacity" or the "primary storage capacity" of cell i,j,k ; the "primary" designation is used to distinguish $SC1_{i,j,k}$ from a secondary storage capacity, which is used when storage term conversion is invoked, as explained in the following section. By using the concept of storage capacity, equation 34 can be rewritten as

$$\frac{\Delta V}{\Delta t} = SC1_{i,j,k} \frac{h_{i,j,k}^m - h_{i,j,k}^{m-1}}{t^m - t^{m-1}} \quad (35)$$

This expression is separated into two terms for inclusion into equation 3:

$$\frac{SC1_{i,j,k} h_{i,j,k}^m}{t^m - t^{m-1}},$$

which is incorporated in the left side of equation 3 through the term $\text{HCOF}_{i,j,k}$, and

$$\frac{-SC1_{i,j,k} h_{i,j,k}^{m-1}}{t^m - t^{m-1}},$$

which is included in the term $\text{RHS}_{i,j,k}$ on the right side of equation 3.

The input to the Layer-Property Flow Package requires specification of specific storage values for each cell of the model if there are one or more transient stress periods. The specific storage values are read layer by layer; and then they are multiplied by the cell areas and layer thickness to create storage capacity values, which are stored in variable $SC1$.

Storage Term Conversion

The primary storage capacity described above, $SC1_{i,j,k}$ is adequate for simulations in which a cell is confined throughout the course of a simulation; however, if the water level is below the top of a cell during a simulation, then the cell is under a water-table condition, and a different storage formulation applies:

$$\frac{\Delta V}{\Delta t} = SY(\text{DELR}_j \text{DEL}C_i) \frac{h_{i,j,k}^m - h_{i,j,k}^{m-1}}{t^m - t^{m-1}} \quad (36)$$

where

SY is the specific yield (dimensionless).

A secondary storage capacity, $SC2_{i,j,k}$, is used to represent specific yield multiplied by cell area, and equation 36 can be written as

$$\frac{\Delta V}{\Delta t} = SC2_{i,j,k} \frac{h_{i,j,k}^m - h_{i,j,k}^{m-1}}{t^m - t^{m-1}} \quad (37)$$

During any one time step, there are four possible storage conditions for each cell:

- The cell is confined for the entire time step,
- The cell is unconfined for the entire time step,
- The cell converts from confined to unconfined, and
- The cell converts from unconfined to confined.

The following expression for rate of accumulation in storage in cell i,j,k is used to handle all four possibilities:

$$\frac{\Delta V}{\Delta t} = \frac{SCB(h_{i,j,k}^m - \text{TOP}_{i,j,k}) + SCA(\text{TOP}_{i,j,k} - h_{i,j,k}^{m-1})}{t^m - t^{m-1}} \quad (38)$$

where

TOP is the elevation of the top of the model cell,

SCA is the storage capacity in effect in cell i,j,k at the start of the time step; and

SCB is the "current" storage capacity—that is, the storage capacity in effect during the current iteration.

Equation 38 handles the four situations as follows: First, consider a case in which the head in cell i,j,k at the beginning of time step m ($h_{i,j,k}^{m-1}$) is above the top of the cell (fig. 9). Because there is no free surface in the cell at the start of the time step, the storage capacity at that time is taken as the confined storage capacity; that is, SCA is set equal to $SC1_{i,j,k}$. If, during a given iteration for time step m , the computed value of head for the end of the time step ($h_{i,j,k}^m$) is found to be above the top of the cell, then SCB for the following iteration is also set equal to $SC1_{i,j,k}$; equation 38 for that iteration then reverts to the form of equation 35, the equation for confined storage. However, if the computed value of $h_{i,j,k}^m$ in a given iteration turns out to be below the top of the cell, as shown in figure 9, the value of SCB for the following iteration is set equal to SC2, the unconfined storage capacity. In this case the computed rate of release of water from storage in the time step has two components:

$$\frac{SC1_{i,j,k}(\text{TOP}_{i,j,k} - h_{i,j,k}^{m-1})}{t^m - t^{m-1}}, \text{ the rate of release from confined or compressive storage, and}$$

$$\frac{SC2_{i,j,k}(h_{i,j,k}^m - \text{TOP}_{i,j,k})}{t^m - t^{m-1}}, \text{ the rate of release from water-table storage.}$$

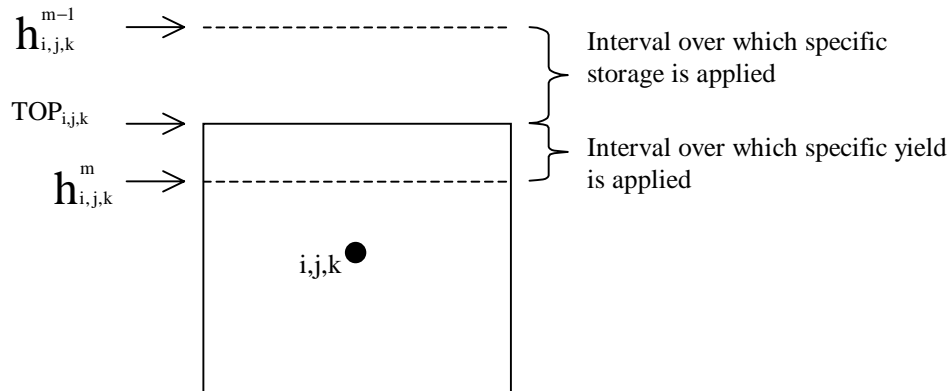


Figure 9.-- A model cell for which two storage factors are used during one time step.

If the head at the beginning of the time step, $h_{i,j,k}$, is below the top of cell i,j,k , so that a free surface exists within the cell, SCA in equation 38 is set equal to $SC2_{i,j,k}$. If, during an iteration for time step m , the computed value of head for the end of the time step is below the top of the cell, SCB in the subsequent iteration is also set equal to $SC2_{i,j,k}$, and equation 38 reverts to the form of equation 37. However, if the computed head for the end of the time step turns out to be above the top of the cell, SCB in the subsequent iteration is set equal to $SC1_{i,j,k}$, the confined storage capacity. This situation occurs during intervals of rising water level, and again two components are computed for the rate of accumulation of water in storage—one corresponding to unconfined or water-table storage and one corresponding to confined or compressive storage.

Equation 38 can be rearranged as follows:

$$\frac{\Delta V}{\Delta t} = \frac{SCB \times h_{i,j,k}^m}{t^m - t^{m-1}} + \frac{SCA(TOP_{i,j,k} - h_{i,j,k}^{m-1}) - SCB \times TOP_{i,j,k}}{t^m - t^{m-1}} \quad (39)$$

Again, $\frac{\Delta V}{\Delta t}$ represents rate of accumulation in storage and as such would appear on the right in equation 2. In the

formulation of equation 3, therefore, the expression $\frac{SCB}{t^m - t^{m-1}}$ is subtracted from $HCOF_{i,j,k}$ on the left-hand side,

while the term

$$\frac{SCA(TOP_{i,j,k} - h_{i,j,k}^{m-1}) - SCB \times TOP_{i,j,k}}{t^m - t^{m-1}}$$

is added to $RHS_{i,j,k}$ on the right-hand side.

For any layer where there is a chance at some point in the simulation that a specific yield should be used rather than a specific storage at one or more cells, then the convertible option should be used for that layer. When this is done, values of specific storage for each cell in the layer are read through the variable Ss , and values of specific yield for each cell in the layer are read through the variable Sy . The specific-storage values are multiplied times cell volume to obtain confined storage capacity, and the specific-yield values are multiplied by

cell area to obtain unconfined storage capacity. Thus, two storage capacities are stored, and the program will use the appropriate value depending on conditions.

Note that unlike the LPF Package, the BCF Package has a special layer type for a water table in layer 1. Equation 37 is then used for calculating storage terms. In LPF, the convertible formulation should be used if layer 1 represents a water table. Also, the elevation of the top of layer 1, which is specified in the discretization file, should be higher than the water level will reach at any time in the simulation. This will result in equation 38 always reverting to equation 37 as described previously. It is the user's responsibility in this situation to check to see that the simulated water level has not exceeded the top elevation, which would trigger the use of confined storage capacity in equation 38. A logical approach for specifying the top elevation for a layer-1 water table is to set it equal to land-surface elevation. That is, in most situations, it is not expected that the water table will exceed land-surface elevation.

Data Requirements

The discretization data—time and space—required by the LPF Package to formulate the flow and storage terms are specified in the Global Process discretization file. All other data required by LPF are read from the LPF data file. There are several optional formulations that are controlled by option variables, which have one value for each layer; that is, the options can be turned on or off on a layer by layer basis. The following option variables are used by LPF:

LAYCBD—Quasi-3D confining bed flag (read as part of the discretization file),
LAYTYP—Layer-type flag,
LAYAVG—Interblock transmissivity flag,
CHANI—Horizontal anisotropy flag and value,
LAYVKA—Vertical anisotropy flag, and
LAYWET—Wetting flag.

The Quasi-3D confining bed flag (LAYCBD) indicates whether a layer is underlain by a confining bed that is simulated using the Quasi-3D Approach. This determines how vertical conductance is calculated.

LAYCBD = 0—there is no Quasi-3D confining bed below the layer, so vertical conductance is calculated using equation 25.

LAYCBD \neq 0—there is a Quasi-3D confining bed below the layer, so vertical conductance is calculated using equation 27.

The layer-type flag (LAYTYP) indicates whether a layer is confined or convertible (between confined and unconfined). This controls whether (1) saturated thickness varies with head, (2) storage term conversion is used, and (3) vertical flow from above is limited under dewatered conditions:

Layer-type flag = 0—The layer is simulated as confined. For a layer of this type, there is no provision for modification of saturated thickness as head varies, for storage term conversion, or for limitation of vertical flow from above if the calculated hydraulic head falls below the top of the cell.

Layer-type flag \neq 0—The layer is simulated as convertible between confined and unconfined. When the calculated hydraulic head is below the top of the cell, all of the options associated with water-table conditions are implemented. Saturated thickness and transmissivity are recalculated at each iteration on the basis of head, and both storage term conversion and limitation of flow from above under dewatered conditions are implemented.

The Interblock transmissivity flag (LAYAVG) indicates the method used for calculating interblock transmissivity:

LAYAVG = 0—Harmonic mean transmissivity is used. This assumes that transmissivity is uniform within a cell, and that changes occur as discrete changes at cell boundaries.

LAYAVG = 1—logarithmic mean transmissivity is used. This assumes that hydraulic conductivity varies linearly between nodes.

LAYAVG = 2—arithmetic-mean thickness and logarithmic-mean hydraulic conductivity are used. This is based on a water-table condition.

The horizontal anisotropy flag (CHANI) indicates whether horizontal anisotropy is constant for a layer, or whether it can vary at each cell in a layer. In either case, it is assumed that the grid is aligned with the principal directions of hydraulic conductivity.

CHANI \leq 0.0—Horizontal anisotropy can vary at each cell in the layer. A separate layer-data variable of the horizontal anisotropy values (HANI) will be read in the input data.

CHANI > 0.0—Horizontal anisotropy is constant for all cells in the layer. The anisotropy is the value of the flag. For example, set CHANI=1.0 for an isotropic layer.

The vertical anisotropy flag (LAYVKA) indicates the type of values that are specified in variable VKA.

LAYVKA = 0—Values of VKA are vertical hydraulic conductivity of the model layer.

LAYVKA \neq 0—Values of VKA are the ratio of horizontal hydraulic conductivity (along rows) to vertical hydraulic conductivity for the model layer.

The wetting flag (LAYWET) indicates whether the wetting capability is on or off.

LAYWET = 0—Wetting capability is off.

LAYWET \neq 0—Wetting capability is on for the layer. This is only valid if the layer-type flag indicates that the layer is convertible (LAYTYP not 0).

Most of the data values that must be specified for the LPF Package are layer data, meaning that the data must be defined for all cells in one or more of the model layers. As used in the above equations that define the LPF formulations, all of the layer-data variables have three-dimensional subscripts, indicating that there are values for all layers; however, some of the variables do not always exist for all layers. The layers for which these layer variables exist are determined by the values in the above option variables. The layer-data variables defined for use by LPF are:

HK—horizontal hydraulic conductivity along rows,

HANI—Ratio of horizontal hydraulic conductivity along columns to hydraulic conductivity along rows,

VKA—Vertical hydraulic conductivity or the ratio of horizontal hydraulic conductivity along rows (HK) to vertical hydraulic conductivity,

VKCB—Vertical hydraulic conductivity of a confining bed below a layer,

Ss—specific storage, and

Sy—specific yield.

WETDRY—wetting threshold combined with a wetting flag.

HK and VKA are always required for all layers. Ss is required for all layers if there are any transient stress periods in the simulation. Sy is required for layers in which LAYTYP \neq 0 and if there are any transient stress periods. The transient/steady-state flags read from the Discretization File determine if storage data are needed.

HANI is required for layers in which $CHANI \leq 0$. VKCB is required for layers in which $LAYCBD \neq 0$. WETDRY is required when $LAYWET \neq 0$.

Defining Layer-Data Variables

The input instructions describe the details of how the LPF input data are read. All of the layer variables can be directly read by using MODFLOW's array readers, and variables HK, HANI, VKA, VKCB, SS, and SY can also be defined by using parameters. The same method of data definition (directly reading or using parameters) must be used for all the layers for a given variable; that is, some of the layers of a particular variable cannot be directly read if some of the layers are defined through parameters and conversely.

When the array readers are used, any of the unused layer data that are included in the input file will cause the data sequence to be misread. Likewise, if any of the required layer arrays are left out of the input file, the data sequence will be misread.

When variables are defined by using array readers, a value must be specified for every cell even if the value will not be used; that is, the array readers read a value for each cell regardless of whether or not the value will be used. For no-flow cells, the values of variables are never used in any calculation, and thus any values may be specified; for constant-head cells, the Ss, Sy, and WETDRY variables are not used but the other variables are.

When variables are defined by using parameters, it is required that a value is defined for variable-head cells ($IBOUND > 0$) and constant-head cells ($IBOUND < 0$). If a value is not assigned at one or more variable- or constant-head cells, then the program will print an error message and stop.

MODFLOW-96 to MODFLOW-2000 Data Conversion Program

MF96TO2K is a utility program that converts MODFLOW-96 input data to MODFLOW-2000 input data. To convert a MODFLOW-96 data set to run with MODFLOW-2000, MF96TO2K reads three files and produces four files: it reads the MODFLOW-96 name, BAS, and BCF files; it produces the discretization file and modified versions of the name, BCF, and BAS files. For confining beds and confined model layers for which top and bottom elevations are not included in the MODFLOW-96 BCF file, MF96TO2K queries the user for these values. Only constant elevation values for each layer can be specified; if non-constant values are required, the user needs to modify the discretization file after it is created by MF96TO2K.

The additional elevation data that MF96TO2K requires in order to create the discretization file are not used by the BCF Package and have no effect on simulation results unless the Horizontal-Flow Barrier Package is being used. The reason these additional values do not affect the BCF Package is that many of the BCF input values directly incorporate the geometry of the system as discussed in the section of this report on the BCF Package. The only elevation data that affect the BCF calculations are the values from the MODFLOW-96 BCF file, which MF96TO2K puts into MODFLOW-2000's discretization file.

The MODFLOW-2000 version of HFB makes use of the additional elevation data that MF96TO2K requests for the discretization file when there are barriers in confined layers. As previously mentioned, if the additional data are not constant for a layer, the user must modify the discretization file after running MF96TO2K. Further, if the Horizontal-Flow Barrier (HFB) Package is used, its input data must be modified, and MF96TO2K does not make this modification.

MF96TO2K functions as follows.

1. It starts by asking for a MODFLOW-96 name file.
2. It then uses this to open the files needed for data conversion.
3. Next, it asks if each layer (except the bottom layer) has a Quasi-3D confining bed under it.
4. It then asks for a constant elevation for any model layers for which elevation data are not included in the MODFLOW-96 BCF file.
5. Finally, MF96TO2K creates new BAS and BCF files using file name extensions BA6 and BC6, respectively. MF96TO2K also creates a discretization file and it overwrites the name file.

After conversion, MODFLOW-2000 should produce the same results as MODFLOW-96 produced.

Adding Packages

As noted in the Introduction section, MODFLOW was originally designed with the goal of making it easy to add new capabilities. The addition of packages is the typical way of enhancing MODFLOW. Packages can be added to the Ground-Water Flow Process of MODFLOW-2000 in much the same way as they could be added to the original versions of MODFLOW. In fact, most packages that have been added to earlier versions of MODFLOW can be added to the Ground-Water Flow Process of MODFLOW-2000 with no change. However, there would probably be an advantage to modifying some packages to include parameters. The most likely cause for an existing package to no longer work in MODFLOW-2000 is the new way in which cell elevations are handled. That is, any package that relies on elevation data as specified in the old BCF Package will require some modification. Another cause for a package to no longer work is the new ability to intermix steady-state and transient stress periods. Examples of packages that require modification are the Transient-leakage (TLK) Package (Leake and others, 1994), the General Finite-Difference Flow (GFD) Package (Harbaugh, 1992), and the Interbed-Storage (IBS) Package (Leake and Prudic, 1991). Examples of packages that require no modification to include in MODFLOW-2000 are the Reservoir (RES) package (Fenske and others, 1996) and the Flow and Head Boundary (FHB) Package (Leake and Lilly, 1997).

Note that packages added to the Ground-Water Flow Process are not necessarily compatible with other processes. There would likely be benefit from adding most hydrologic packages to other processes within MODFLOW-2000, but this is beyond the scope of this report. The documentation for new packages should include information about the processes that they work with. Previously developed packages must be evaluated to determine compatibility with other processes, and the documentation of new processes should describe the degree of compatibility with existing packages to the extent possible.

INPUT INSTRUCTIONS

Input instructions for MODFLOW-2000 are divided into four major sections: Global Process, Ground-Water Flow Process, Array Utility Modules, and List Utility Module. The Global Process reads several files, and each file is described in a separate subsection. The Ground-Water Flow Process reads at least one file for each package, and these files are also described separately. The instructions for a file are grouped by numbered items, and each item consists of input variables. If a variable is represented in the program by a variable of the same name, then the variable is shown in all upper case. Variables that are not actual program variables are capitalized. Each variable is defined after all the items are listed.

An input variable may include a single value or multiple values. One-dimensional variables are multi-valued variables in which the number of values is indicated by a single number in parentheses after the name. Two-dimensional variables are multi-valued variables in which the number of values is indicated by two numbers in parentheses after the name. The total number of values represented is the product of the two numbers. A layer variable is a two-dimensional variable that includes one value for each cell in a layer. Therefore, the column and row dimensions are the numbers in parentheses after the variable. For example, "IBOUND(NCOL,NROW)" indicates that IBOUND is a layer variable. Most multi-value variables are read by utility modules, and if so, the utility module (U1DREL, U2DREL, or U2DINT) is named along with the variable. Module U1DREL is used for reading one-dimensional variables that are real numbers, U2DREL is used for reading two-dimensional variables that are real numbers, and U2DINT is used for reading two-dimensional variables that are integer numbers.

Some items consist of several variables, and the item can be repeated multiple times. This kind of item is called a list item. List items are usually read by the list utility module, ULSTRD. The use of ULSTRD is mentioned as part of the note that describes the repeated item.

The input data for each item must start on a new record. (A record is a line in a file.) It is assumed that all variables for an item will be contained in a single record unless the item consists of a multi-valued variable. That is, a multi-valued variable may occupy multiple records. When reading a layer variable, the data for each row must start on a new record.

Each input variable has a data type, which can be Real, Integer, or Character. Integers are whole numbers, and these values must not include a decimal point or exponent. Real numbers can include a decimal point and an exponent. If no decimal point is included in the entered value, then the decimal point is assumed to be at the right side of the value. Any printable character is allowed for character variables. All variables starting with the letters I-N are integers. Variables starting with the letters A-H and O-Z are either real numbers or character data, and most of these variables are real numbers; that is, there are few character variables in MODFLOW. It should be assumed that a variable starting with A-H or O-Z is a real number unless otherwise stated in the variable definitions. This data-type convention is used for all input variables regardless of whether a variable is an actual program variable.

Both free and fixed formatting are used as specified throughout the input instructions. With fixed format, a data value has a specified column location, called the field, within a record. If a numeric value does not require the entire field width, the value should be right justified within the field, and blanks should fill the remaining width of the field. The justification of character data within a field does not matter unless specifically noted.

Free format is similar to Fortran's list directed input. With free format, values are not required to occupy a fixed number of columns in a record. Each value can occupy one or more columns as required to represent it; however, the values must still be included in the prescribed order. One or more spaces, or a single comma optionally combined with spaces, must separate adjacent values. Also, a numeric value of zero must be explicitly represented with 0 and not by one or more spaces when free format is used, because there is no way to detect the difference between a space that represents 0 and a space that represents a value separator.

Two capabilities included in Fortran's list-directed input are not included in the free-format input implemented in MODFLOW-2000.

1. Null values in which input values are left unchanged from their previous values are not allowed. In general, MODFLOW's input values are not defined prior to their input.
2. A "/" cannot be used to terminate an input record without including values for all the variables; data values for all input variables must be explicitly specified on an input record.

For character data, MODFLOW's free format implementation is less stringent than the list-directed input of Fortran 77. Fortran requires character data to be delineated by apostrophes. MODFLOW does not require apostrophes unless a blank or a comma is part of a character variable.

An example problem is included in the Example section. That section shows input files that illustrate the use of parameters.

Running MODFLOW-2000

MODFLOW-2000 is executed in the same manner as MODFLOW-96. Except as described in the following paragraphs, when MODFLOW-2000 is started, it prompts the user to enter the name of the name file, which is described below. Once the name is entered, MODFLOW-2000 runs without further user interaction.

Provision has been made to allow a series of simulations to be run without the need to manually restart MODFLOW-2000 for each simulation and respond to the prompt for the name file. This is called batch mode and is done through the use of a special file that must be named `modflow.bf`. On case-sensitive operating systems, such as Unix, lower case letters are needed. If a file with this name exists in the user's current directory when MODFLOW-2000 is invoked, the user is not prompted for a name file. Instead, MODFLOW-2000 reads the names of name files from `modflow.bf`. Each line of the `modflow.bf` file contains the name of a single name file, and each name can be up to 80 characters long.

When batch mode is used, a report file, named `modbatch.rpt`, is created. Before starting a simulation, the name of the name file is written to the report file. If the name file exists, the simulation is executed; when done, another message is written to the report file. If a name file does not exist, a message is written to the report file and processing continues with the next name file in `modflow.bf`.

When batch mode is used to run MODFLOW-2000, two additional file units are used—97 and 98. These units cannot be used by any of the MODFLOW-2000 simulations. Within the name files listed in `modflow.bf`, the

file units listed in one name file can be the same as those used in any of the other name files because each simulation is executed separately.

Global Process Input Instructions

Name File

The name file contains the names of most input and output files used in a model simulation and controls the parts of the model program that are active. (“OPEN/CLOSE” files, described in the Input Instructions for Array Reading Utility Modules section, are not included in the name file.) The name file is read on unit 99. The name file is constructed as follows.

FOR EACH SIMULATION

1. Ftype Nunit Fname

The name file contains one of the above records (item 1) for each file. All variables are free format. The length of each record must be 199 characters or less. The records can be in any order except for records where Ftype (file type) is “GLOBAL” or “LIST” as described below.

Comment records are indicated by the # character in column one and can be located anywhere in the file. Any text characters can follow the # character. Comment records have no effect on the simulation; their purpose is to allow users to provide documentation about a particular simulation. All comment records after the first item-1 record are written in the listing file.

Explanation of Variables in the Name File

Ftype—is the file type, which must be one of the following character values. Ftype may be entered in all uppercase, all lowercase, or any combination.

GLOBAL for the global listing file—if this type is not present, the LIST file is used for the global listing file as well as for the forward run listing file.

LIST for the forward run listing file—if this type is not present, the GLOBAL file is used for the forward run listing file as well as for the global listing file.

The name file must always include a record that specifies GLOBAL or LIST for Ftype. Both records can be included, and if so, the GLOBAL record must be the first non-comment record and the LIST record must be the second non-comment record. If only one of the records is included, it must be the first non-comment record.

DIS for the discretization file

MULT for the multiplier array file

ZONE for the zone array

BAS6 for the Ground-Water Flow Process Basic Package

OC for the Ground-Water Flow Process Output Control Option

BCF6 for the Ground-Water Flow Process Block-Centered Flow Package

LPF for the Ground-Water Flow Process Layer Property Flow package

HFB6 for the Ground-Water Flow Process Horizontal Flow Barrier Package

RCH for the Ground-Water Flow Process Recharge Package

RIV for the Ground-Water Flow Process River Package

WEL for the Ground-Water Flow Process Well Package

DRN for the Ground-Water Flow Process Drain Package
GHB for the Ground-Water Flow Process General-Head Boundary Package
EVT for the Ground-Water Flow Process Evapotranspiration Package
CHD for the Ground-Water Flow Process Constant-Head Package

SIP for the Strongly Implicit Procedure Package
SOR for the Slice-Successive Over-Relaxation Package
PCG for the Preconditioned Conjugate-Gradient Package
DE4 for the Direct Solution Package

DATA(BINARY) for binary (unformatted) files, such as those used to save cell-by-cell budget data and binary (unformatted) head and drawdown data. Files of this type are rewound at the start of each parameter-estimation iteration.

DATA for formatted (text) files, such as those used to save formatted head and drawdown and for input of data from files that are separate from the primary package input files. Files of this type are rewound at the start of each parameter-estimation iteration.

DATAGLO(BINARY) for binary (unformatted) files, such as those used to save cell-by-cell budget data and binary (unformatted) head and drawdown data. Files of this type are not rewound at the start of each parameter-estimation iteration; that is, output from each successive parameter iteration accumulates rather than overwriting information from the previous iteration.

DATAGLO for formatted (text) files, such as those used to save formatted head and drawdown and for input of data from files that are separate from the primary package input files. Files of this type are not rewound at the start of each parameter-estimation iteration; that is, output from each successive parameter iteration accumulates rather than overwriting information from the previous iteration.

Nunit—is the Fortran unit to be used when reading from or writing to the file. Any legal unit number on the computer being used can be specified except units 96-99. Unit 99 is used for the name file and for reading multi-valued variables using the OPEN/CLOSE option of the utility modules (see Input Instructions for Array Reading Utility Modules section). Units 97 and 98 are used for batch files as explained below. Unit 96 is used for a file to which error messages may be written when the Sensitivity Process is active. The unit number for each file must be unique.

Fname—is the name of the file, which is a character value. Pathnames may be specified as part of Fname.

Discretization File

Discretization information is read from the file that is specified by "DIS" as the file type.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. NLAY NROW NCOL NPER ITMUNI LENUNI
2. LAYCBD(NLAY)
3. DELR(NCOL) - U1DREL
4. DELC(NROW) - U1DREL
5. Top(NCOL,NROW) - U2DREL
6. BOTM(NCOL,NROW) - U2DREL

Item 6 is repeated for each model layer and Quasi-3D confining bed in the grid. These layer variables are read in sequence going down from the top of the system. Thus, the number of BOTM arrays must be NLAY plus the number of Quasi-3D confining beds.

FOR EACH STRESS PERIOD

7. PERLEN NSTP TSMULT Ss/tr

Explanation of Variables Read from the Discretization File

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NLAY—is the number of layers in the model grid.

NROW—is the number of rows in the model grid.

NCOL—is the number of columns in the model grid.

NPER—is the number of stress periods in the simulation.

ITMUNI—indicates the time unit of model data, which must be consistent for all data values that involve time. For example, if years is the chosen time unit, stress-period length, time-step length, transmissivity, and so on, must all be expressed using years for their time units. Note that the program will still run even if “undefined” time units are specified because the fundamental equations used in MODFLOW do not require that the time unit be identified. But be sure to use consistent units for all input data even when ITMUNI indicates an undefined time unit. When the time unit is defined, MODFLOW uses it to print a table of elapsed simulation time:

0 - undefined	3 - hours
1 - seconds	4 - days
2 - minutes	5 - years

LENUNI—indicates the length unit of model data, which must be consistent for all data values that involve length. For example, if feet is the chosen length unit, grid spacing, head, hydraulic conductivity, water volumes, and so forth, must all be expressed using feet for their length units. Note that the program will still run even if “undefined” length units are specified because the fundamental equations used in MODFLOW do not require that the length unit be identified. But be sure to use consistent units for all input data even when LENUNI indicates an undefined length unit:

0 - undefined
1 - feet

- 2 - meters
- 3 - centimeters

LAYCBD—is a flag, with one value for each model layer, that indicates whether or not a layer has a Quasi-3D confining bed below it. 0 indicates no confining bed, and not zero indicates a confining bed. LAYCBD for the bottom layer must be 0.

DELR—is the cell width along rows. Read one value for each of the NCOL columns. This is a multi-value one-dimensional variable with one value for each column.

DELC—is the cell width along columns. Read one value for each of the NROW rows. This is a multi-value one-dimensional variable with one value for each row.

Top—is the top elevation of layer 1. For the common situation in which the top layer represents a water-table aquifer, it may be reasonable to set Top equal to land-surface elevation.

BOTM—is the bottom elevation of a model layer or a Quasi-3d confining bed.

PERLEN—is the length of a stress period.

NSTP—is the number of time steps in a stress period.

TSMULT—is the multiplier for the length of successive time steps. The length of a time step is calculated by multiplying the length of the previous time step by TSMULT. The length of the first time step, Δt_1 , is related to PERLEN, NSTP, and TSMULT by the relation

$$\Delta t_1 = \text{PERLEN} \left(\frac{\text{TSMULT} - 1}{\text{TSMULT}^{\text{NSTP}} - 1} \right) .$$

Ss/tr—is a character variable that indicates if the stress period is transient or steady state. The only allowed options are “SS” and “TR”, but these are case insensitive.

Multiplier File

Input to define multiplier arrays is read from the file that is specified with "MULT" as the file type. Multiplier arrays can be used to calculate layer variables from parameter values.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. NML

2. MLTNAM [*FUNCTION*]

If Item 2 does not contain the optional *FUNCTION* keyword, read Item 3:

3. [RMLT (NCOL , NROW)] - U2DREL

Otherwise, if Item 2 contains the optional *FUNCTION* keyword, read Item 4:

4. [MLTNAM1 [op1 MLTNAM2] [op2 MLTNAM3] [op3 MLTNAM4] . . .]

Repeat items 2-4 for each of the NML multiplier arrays.

Explanation of Variables Read from the Multiplier File

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NML—is the number of multiplier arrays to be defined.

MLTNAM—is the name of a multiplier array. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case are equivalent. The name “NONE” should not be used for a multiplier array because it is a reserved word. The optional keyword, *FUNCTION*, indicates that the multiplier array will be constructed from other multiplier arrays that have already been defined. Construction is by arithmetic combinations of the multipliers; see the explanation below for variable op1, op2, op3

RMLT—is a multiplier array.

MLTNAM1, MLTNAM2, MLTNAM3, ...—are the names of multiplier arrays that have already been defined.

op1, op2, op3, ...—are arithmetic operators used to define a multiplier array based on other multiplier arrays. Each operator can be either “+”, “-”, “*”, or “/”. Operations are applied from left to right to each array element. The operators must be separated from the multiplier array names by at least one space.

Example Multiplier Array Input Using the FUNCTION Keyword

The following example input illustrates the use of the *FUNCTION* keyword to construct a multiplier array from other multiplier arrays. In this example, three multiplier arrays are defined. The first two arrays (named M1 and M2) are read using the U2DREL utility array reader (item 3), and the third array (named M3) is defined as the sum of the other two. In this example, a model layer has 5 rows and 4 columns.

3

M1

INTERNAL	1.0	(4F6.0)	0
1.0	1.1	1.2	1.3
1.0	1.1	1.2	1.3
2.0	2.2	2.4	2.6
2.0	2.2	2.4	2.6
1.0	1.1	1.2	1.3

M2

INTERNAL	1.0	(4F6.0)	0
5.0	5.1	5.2	5.3
5.0	5.1	5.2	5.3
6.0	6.1	6.2	6.3
6.0	6.1	6.2	6.3
5.0	5.1	5.2	5.3

M3 FUNCTION

M1 + M2

The resulting values for multiplier M3 are:

6.0	6.2	6.4	6.6
6.0	6.2	6.4	6.6
8.0	8.3	8.6	8.9
8.0	8.3	8.6	8.9
6.0	6.2	6.4	6.6

Zone File

Input to define zone arrays is read from the file that is specified with "ZONE" as the file type. Zone arrays can be used to specify the cells in a layer variable that are associated with a parameter.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. NZN

2. ZONNAM

3. IZON(NCOL, NROW) - U2DINT

Items 2-3 are repeated for each of the NZN zone arrays.

Explanation of Variables Read from the Zone File

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NZN—is the number of zone arrays to be defined.

ZONNAM—is the name of a zone array. This name can consist of 1 to 10 characters and is not case sensitive; that is, any combination of the same characters with different case are equivalent. The name “ALL” should not be used for a zone array because it is a reserved word.

IZON—is a zone array.

Ground-Water Flow Process Input Instructions

Basic Package

Input for the Ground-Water Flow Process Basic (BAS) Package except for output control input is read from the file that is specified with “BAS6” as the file type. The file type has been changed from that used in MODFLOW-96, which was “BAS”, because the input data are not compatible.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—“#” must be in column 1. Item 0 can be repeated multiple times.
1. Options (199 text characters)
If there are no options to specify, then a blank record must be included for Item 1.
2. IBOUND(NCOL,NROW) or (NCOL,NLAY) -- U2DINT
If not a cross section, a layer variable is read for each layer in the grid.
If a cross section, NLAY rows of NCOL values are read.
3. HNOFLO (10-character field unless Item 1 contains 'FREE'.)
4. STRT(NCOL,NROW) or (NCOL,NLAY) -- U2DREL
If not a cross section, a layer variable is read for each layer in the grid.
If a cross section, NLAY rows of NCOL values are read.

Explanation of Variables Read by the BAS Package

Text—is a character variable that starts in column 2. The first two comment lines will become variable HEADNG, which is used as a printout title throughout the program. (If there are no comment lines, then HEADNG will be blank.) HEADNG is limited to 80 columns, but subsequent Text lines can be up to 199 columns. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

Options—is a character variable that is scanned for words (separated by one or more spaces) that specify program options. Three options are currently recognized. Unrecognized words are ignored, and a word may be specified in either uppercase or lowercase. A blank record is acceptable and indicates no options.

XSECTION indicates that the model is a 1-row cross section for which STRT and IBOUND should each be read as single two-dimensional variables with dimensions of NCOL and NLAY. Likewise, head and drawdown should be printed and saved in disk files as single two-dimensional variables.

CHTOCH indicates that flow between adjacent constant-head cells should be calculated.

FREE indicates that free format is used for input variables throughout the Basic Package and other packages as indicated in their input instructions. Be sure that all variables read using free format have a non-blank value and that a comma or at least one blank separates all adjacent values.

IBOUND—is the boundary variable. One value is read for every model cell. Usually, these values are read a layer at a time; however, when the XSECTION option is specified, a single variable for the cross section is read. Note

that although IBOUND is read as one or more two-dimensional variables, it is stored internally as a three-dimensional variable.

If $IBOUND(J,I,K) < 0$, cell J,I,K has a constant head.

If $IBOUND(J,I,K) = 0$, cell J,I,K is inactive.

If $IBOUND(J,I,K) > 0$, cell J,I,K is active.

HNOFLO—is the value of head to be assigned to all inactive (no flow) cells ($IBOUND = 0$) throughout the simulation. Because head at inactive cells is unused in model calculations, this does not affect model results but serves to identify inactive cells when head is printed. This value is also used as drawdown at inactive cells if the drawdown option is used. Even if the user does not anticipate having inactive cells, a value for HNOFLO must be entered.

STRT—is initial (starting) head—that is, head at the beginning of the simulation. STRT must be specified for all simulations, including steady-state simulations. One value is read for every model cell. Usually, these values are read a layer at a time. When the XSECTION option is specified, however, a single array for the cross section is read. For simulations in which the first stress period is steady state, the values used for STRT generally do not affect the simulation (exceptions may occur if cells go dry and(or) rewet). The execution time, however, will be less if STRT includes hydraulic heads that are close to the steady-state solution.

Output Control Option

Input to the Output Control Option of the Ground-Water Flow Process is read from the file that is specified as type "OC" in the name file. If no "OC" file is specified, default output control is used. Under the default, head and overall budget are written to the listing file (printed) at the end of every stress period. The default printout format for head and drawdown is 10G11.4.

There are two ways to specify Output Control data: words or numeric codes. One of these methods must be used throughout any simulation.

Output Control Using Words

Recognized words are shown in bold italics; these words must be entered exactly as shown except that they may be entered in either uppercase or lowercase. Optional parts of records are shown in brackets. One or more spaces must separate each word or variable, and the total record length must not exceed 199 characters.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. Any combination of the following records:

HEAD PRINT FORMAT IHEDFM
HEAD SAVE FORMAT CHEDFM [**LABEL**] Omit to obtain an unformatted file.
HEAD SAVE UNIT IHEDUN

DRAWDOWN PRINT FORMAT IDDNFM
DRAWDOWN SAVE FORMAT CDDNFM [**LABEL**] Omit to obtain an unformatted file.
DRAWDOWN SAVE UNIT IDDNUN

IBOUND SAVE FORMAT CBOUFM [**LABEL**]
IBOUND SAVE UNIT IBOUUN

COMPACT BUDGET [**AUX** or **AUXILIARY**]

If the final "**COMPACT BUDGET**" option is used, cell-by-cell budget files produced require less disk space than files written by MODFLOW-88; however, programs that read these data using the specifications of MODFLOW-88 will need to be modified. If this option is not used, MODFLOW-2000 will write the files using the MODFLOW-88 method. The optional word "**AUX**" (or "**AUXILIARY**") after "**COMPACT BUDGET**" indicates that auxiliary data that are defined in packages (see input data for the RIV, WEL, DRN, and GHB Packages) should be saved in the budget file along with budget data.

PRINT indicates that the model output is written to the LIST file.

SAVE indicates that output is written to the indicated file unit. Drawdown and head are saved as text files if the **SAVE FORMAT** record is included. If the **SAVE FORMAT** record is not used, then the file is written as a binary (unformatted) file. Binary files are usually more compact than text files, but they are not generally transportable among different computer operating systems or different Fortran compilers.

FOR EACH TIME STEP FOR WHICH OUTPUT IS DESIRED

2. **PERIOD** IPEROC **STEP** ITSOC

3. Any combination of the following records:

PRINT HEAD [list layers if all layers not desired]
PRINT DRAWDOWN [list layers if all layers not desired]

PRINT BUDGET Write overall volumetric budget to the LIST file.
SAVE HEAD [list layers if all layers not desired]
SAVE DRAWDOWN [list layers if all layers not desired]
SAVE IBOUND [list layers if all layers not desired]
SAVE BUDGET Write cell-by-cell budget data to a file.

Item 2 and one or more item-3 records are specified for each time for which output is desired. These records must be in the order of increasing simulation time.

Explanation of Variables Read by Output Control Using Words

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

IHEDFM—is a code for the format in which heads will be printed. (Positive values for wrap format; negative values for strip format.)

0 - 10G11.4	11 - 20F5.4
1 - 11G10.3	12 - 10G11.4
2 - 9G13.6	13 - 10F6.0
3 - 15F7.1	14 - 10F6.1
4 - 15F7.2	15 - 10F6.2
5 - 15F7.3	16 - 10F6.3
6 - 15F7.4	17 - 10F6.4
7 - 20F5.0	18 - 10F6.5
8 - 20F5.1	19 - 5G12.5
9 - 20F5.2	20 - 6G11.4
10 - 20F5.3	21 - 7G9.2

CHEDFM—is a character value that specifies the format for saving heads, and can only be specified if the word method of output control is used. The format must contain 20 characters or less and must be a valid Fortran format that is enclosed in parentheses. The format must be enclosed in apostrophes if it contains one or more blanks or commas. The optional word **LABEL** after the format is used to indicate that each layer of output should be preceded with a line that defines the output (simulation time, the layer being output, and so forth). If there is no record specifying CHEDFM, then heads are written to a binary (unformatted) file. Binary files are usually more compact than text files, but they are not generally transportable among different computer operating systems or different Fortran compilers.

IHEDUN—is the unit number on which heads will be saved.

IDDNFM—is a code for the format in which drawdowns will be printed. The codes are the same as for IHEDFM.

CDDNFM—is a character value that specifies the format for saving drawdown, and can only be specified if the word method of output control is used. The format must contain 20 characters or less and must be a valid Fortran format that is enclosed in parentheses. The format must be enclosed in apostrophes if it contains one or more blanks or commas. The optional word **LABEL** after the format is used to indicate that each layer of output should be preceded with a line that defines the output (simulation time, the layer being output, and so forth). If there is no record specifying CDDNFM, then drawdown is written to a binary (unformatted) file. Binary files are usually more compact than text files, but they are not generally transportable among different computer operating systems or different Fortran compilers.

IDDNUN—is the unit number on which drawdowns will be saved.

CBOUFM—is a character value that specifies the format for saving IBOUND, and can only be specified if the word method of output control is used. The format must contain 20 characters or less and must be a valid Fortran format that is enclosed in parentheses. The format must be enclosed in apostrophes if it contains one or more blanks or commas. The optional word **LABEL** is used to indicate that each layer of output should be preceded with a line that defines the output (simulation time, the layer being output, and so forth). If there is no record

specifying CBOUFM, then IBOUND is written using format (20I4). IBOUND is never written as a binary (unformatted) file.

IBOUUN—is the unit number on which drawdowns will be saved.

IPEROC—is the stress period number at which output is desired.

ITSOC—is the time step number (within a stress period) at which output is desired.

Example Output Control Input Using Words

The first line cannot be blank, but after the first line blank lines are ignored when the word method is used to specify Output Control data. Indented lines are allowed because of the use of free format input.

```
HEAD PRINT FORMAT 15
HEAD SAVE FORMAT (20F10.3) LABEL
HEAD SAVE UNIT 30
COMPACT BUDGET FILES
DRAWDOWN PRINT FORMAT 14
```

```
PERIOD 1 STEP 1
  PRINT HEAD 2 6
  PRINT DRAWDOWN
  PRINT BUDGET
  SAVE BUDGET
  SAVE HEAD
```

```
PERIOD 1 STEP 7
  SAVE HEAD 1 3 5
  PRINT DRAWDOWN
  SAVE BUDGET
```

```
PERIOD 2 STEP 5
  PRINT HEAD
  PRINT BUDGET
  SAVE BUDGET
  SAVE HEAD
```

Output Control Using Numeric Codes

All variables are free format if the word FREE is specified in Item 1 of the Basic Package input file; otherwise, the variables all have 10-character fields.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—“#” must be in column 1. Item 0 can be repeated multiple times.

```
1.   IHEDFM      IDDNFM      IHEDUN      IDDNUN
```

FOR EACH TIME STEP

```
2.   INCODE      IHDDFL      IBUDFL      ICBCFL
3.   Hdpr        Ddpr        Hdsv        Ddsv
```

(Item 3 is read 0, 1, or NLAY times, depending on the value of INCODE.)

Explanation of Variables Read by Output Control Using Numeric Codes

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

IHEDFM—is a code for the format in which heads will be printed. See the description above in the explanation of variables read by output control using words.

IHEDUN—is the unit number on which heads will be saved.

IDDNFM—is a code for the format in which drawdowns will be printed. The codes are the same as for IHEDFM.

IDDNUN—is the unit number on which drawdowns will be saved.

INCODE—is the code for reading Item 3.

If $INCODE < 0$, Item 3 flags are used from the last time step. Item 3 is not read.

If $INCODE = 0$, all layers are treated the same way. Item 3 will consist of one record.

If $INCODE > 0$, Item 3 will consist of one record for each layer.

IHDDFL—is a head and drawdown output flag. This flag allows Item 3 flags to be specified in an early time step and then used or not used in subsequent time steps. Thus, it may be possible to use IHDDFL to avoid resetting Item 3 flags every time step.

If $IHDDFL = 0$, no heads or drawdowns will be printed or saved regardless of which Item 3 flags are specified.

If $IHDDFL \neq 0$, heads and drawdowns will be printed or saved according to the Item 3 flags.

IBUDFL—is a budget print flag.

If $IBUDFL = 0$, overall volumetric budget will not be printed.

If $IBUDFL \neq 0$, overall volumetric budget will be printed.

ICBCFL—is a flag for writing cell-by-cell flow data.

If $ICBCFL = 0$, cell-by-cell flow terms are not written to any file.

If $ICBCFL \neq 0$, cell-by-cell flow terms are written to the LIST file or a budget file depending on flags set in the component of flow packages, that is, IWELCB, IRCHCB, and so forth.

Hdpr—is the output flag for head printout.

If $Hdpr = 0$, head is not printed for the corresponding layer.

If $Hdpr \neq 0$, head is printed for the corresponding layer.

Ddpr—is the output flag for drawdown printout.

If $Ddpr = 0$, drawdown is not printed for the corresponding layer.

If $Ddpr \neq 0$, drawdown is printed for the corresponding layer.

Hdsv—is the output flag for head save.

If $Hdsv = 0$, head is not saved for the corresponding layer.

If $Hdsv \neq 0$, head is saved for the corresponding layer.

Ddsv—is the output flag for drawdown save.

If $Ddsv = 0$, drawdown is not saved for the corresponding layer.

If $Ddsv \neq 0$, drawdown is saved for the corresponding layer.

Block-Centered Flow Package

This version of the Block-Centered Flow (BCF) Package combines the capabilities documented in McDonald and Harbaugh (1988), McDonald and others (1992), and Goode and Appel (1992). Input for the BCF Package is read from the file that is type "BCF6" in the name file. The file type has been changed from that used in MODFLOW-96, which was "BCF", because the input data are not compatible.

FOR EACH SIMULATION

1. IBCFCB HDRY IWDFLG WETFCT IWETIT IHDWET
These 6 variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the variables all have 10-character fields.
2. Ltype(NLAY)
Read one value for each layer. These values are free format if the word FREE is specified in Item 1 of the Basic Package input file; otherwise, they have fixed-format fields. The fixed format fields are each 2 characters wide with 40 values per line. Use only as many lines as required for the number of model layers.
3. TRPY(NLAY) -- U1DREL

A subset of the following two-dimensional variables are used to describe each layer. The variables needed for each layer depend on the layer-type code (LAYCON, which is defined as part of the Item-2 Ltype), whether the simulation has any transient stress periods (at least one stress period defined in the Discretization File specifies Ss/Tr as "TR"), and if the wetting capability is active (IWDFLG not 0). If a variable is not needed, it must be omitted. In no situation will all variables be required. The required variables (Items 4-9) for layer 1 are read first; then the variables for layer 2 and so forth.

4. [Sf1(NCOL,NROW)] -- U2DREL If there is at least one transient stress period.

If LAYCON is 0 or 2 (see Ltype), then read item 5.

5. [Tran(NCOL,NROW)] -- U2DREL.

Otherwise, if LAYCON is 1 or 3 (see Ltype), read item 6.

6. [HY(NCOL,NROW)] -- U2DREL
7. [Vcont(NCOL,NROW)] -- U2DREL If not the bottom layer.
8. [Sf2(NCOL,NROW)] -- U2DREL If there is at least one transient stress period and LAYCON (see Ltype) is 2 or 3.
9. [WETDRY(NCOL,NROW)] -- U2DREL If IWDFLG is not 0 and LAYCON is 1 or 3 (see Ltype).

Explanation of Variables Read by the BCF Package

IBCFCB—is a flag and a unit number.

If IBCFCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control. The terms that are saved are storage, constant-head flow, and flow between adjacent cells.

If IBCFCB = 0, cell-by-cell flow terms will not be written.

If IBCFCB < 0, cell-by-cell flow for constant-head cells will be written in the listing file when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control. Cell-by-cell flow to storage and between adjacent cells will not be written to any file.

HDRY—is the head that is assigned to cells that are converted to dry during a simulation. Although this value plays no role in the model calculations, it is useful as an indicator when looking at the resulting heads that are output from the model. HDRY is thus similar to HNOFLO in the Basic Package, which is the value assigned to cells that are no-flow cells at the start of a model simulation.

IWDFLG—is a flag that determines if the wetting capability is active.

If IWDFLG = 0, the wetting capability is inactive.

If IWDFLG is not 0, the wetting capability is active.

WETFCT—is a factor that is included in the calculation of the head that is initially established at a cell when it is converted from dry to wet. (See IHDWET.)

IWETIT—is the iteration interval for attempting to wet cells. Wetting is attempted every IWETIT iteration. If using the PCG solver (Hill, 1990), this applies to outer iterations, not inner iterations. If IWETIT is 0, it is changed to 1.

IHDWET—is a flag that determines which equation is used to define the initial head at cells that become wet:

If IHDWET = 0, equation 33A (equation 3a in McDonald and others, 1992) is used:

$$h = BOT + WETFCT (h_n - BOT)$$

If IHDWET is not 0, equation 33B (equation 3b in McDonald and others, 1992) is used:

$$h = BOT + WETFCT (WETDRY)$$

Ltype—contains a combined code for each layer that specifies both the layer type (LAYCON) and the method of computing interblock conductance. Use as many records as needed to enter a value for each layer. Values are two-digit numbers:

The left digit defines the method of calculating interblock transmissivity. The methods are described by Goode and Appel (1992).

0 or blank—harmonic mean (the method used in MODFLOW-88).

1—arithmetic mean

2—logarithmic mean

3—arithmetic mean of saturated thickness and logarithmic-mean hydraulic conductivity.

The right digit defines the layer type (LAYCON), which is the same as in MODFLOW-88:

0—confined—Transmissivity and storage coefficient of the layer are constant for the entire simulation.

1—unconfined—Transmissivity of the layer varies. It is calculated from the saturated thickness and hydraulic conductivity. The storage coefficient is constant. This type code is valid only for layer 1.

2—confined/unconfined—Transmissivity of the layer is constant. The storage coefficient may alternate between confined and unconfined values. Vertical flow from above is limited if the layer desaturates.

3—confined/unconfined—Transmissivity of the layer varies. It is calculated from the saturated thickness and hydraulic conductivity. The storage coefficient may alternate between confined and unconfined values. Vertical flow from above is limited if the aquifer desaturates.

TRPY—is a one-dimensional variable containing a horizontal anisotropy factor for each layer. It is the ratio of transmissivity or hydraulic conductivity (whichever is being used) along a column to transmissivity or hydraulic conductivity along a row. Set to 1.0 for isotropic conditions. This is a single variable with one value per layer. Do not read a variable for each layer—that is, include only one array control record for the entire variable. Note that the LPF Package can be used to represent horizontal anisotropy that varies in magnitude within a model layer.

Sf1—is the primary storage coefficient. Read only if there are one or more transient stress periods specified in the Discretization File. For LAYCON equal to 1, Sf1 will always be specific yield, whereas for LAYCON equal to 2 or 3, Sf1 will always be confined storage coefficient. For LAYCON equal to 0, Sf1 would normally be confined storage coefficient; however, a LAYCON value of 0 can also be used to simulate water-table conditions where drawdowns are expected to remain everywhere a small fraction of the saturated thickness, and where there is no layer above, or flow from above is negligible. In this case, specific yield values would be entered for Sf1.

Tran—is the transmissivity along rows. Tran is multiplied by TRPY to obtain transmissivity along columns. Read only for layers where LAYCON is 0 or 2.

HY—is the hydraulic conductivity along rows. HY is multiplied by TRPY to obtain hydraulic conductivity along columns. Read only for layers where LAYCON is 1 or 3.

Vcont—is the vertical hydraulic conductivity divided by the thickness from a layer to the layer below. The value for a cell is the hydraulic conductivity divided by thickness for the material between the node in that cell and the node in the cell below. Because there is not a layer beneath the bottom layer, Vcont cannot be specified for the bottom layer.

Sf2—is the secondary storage coefficient. Read only for layers where LAYCON is 2 or 3 and only if there are one or more transient stress periods specified in the Discretization File. The secondary storage coefficient is always specific yield.

WETDRY—is a combination of the wetting threshold and a flag to indicate which neighboring cells can cause a cell to become wet. If $WETDRY < 0$, only the cell below a dry cell can cause the cell to become wet. If $WETDRY > 0$, the cell below a dry cell and the four horizontally adjacent cells can cause a cell to become wet. If $WETDRY$ is 0, the cell cannot be wetted. The absolute value of $WETDRY$ is the wetting threshold. When the sum of BOT and the absolute value of $WETDRY$ at a dry cell is equaled or exceeded by the head at an adjacent cell, the cell is wetted. Read only if LAYCON is 1 or 3 and IWDFLG is not 0.

Layer-Property Flow Package

Input for the Layer-Property Flow (LPF) Package is read from the file that is type "LPF" in the name file. Free format is used for reading all values. The LPF Package is an alternative to the BCF Package. Both packages should not be used simultaneously.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. ILPFCB HDRY NPLPF
2. LAYTYP (NLAY)
3. LAYAVG (NLAY)
4. CHANI (NLAY)
5. LAYVKA (NLAY)
6. LAYWET (NLAY)
7. [WETFCT IWETIT IHDWET]
(Include item 7 only if LAYWET indicates at least one wettable layer.)
8. [PARNAM PARTYP Parval NCLU]
9. [Layer Mltarr Zonarr IZ]
Each repetition of Item 9 is called a parameter cluster. Repeat Item 9 NCLU times.
Repeat Items 8-9 for each parameter to be defined (that is, NPLPF times).

A subset of the following two-dimensional variables is used to describe each layer. All the variables that apply to layer 1 are read first, followed by layer 2, followed by layer 3, and so forth. If a variable is not required due to simulation options (for example, Ss and Sy for a completely steady-state simulation), then it must be omitted from the input file.

These variables are either read by the array-reading utility module, U2DREL, or they are defined through parameters. If a variable is defined through parameters, then the variable itself is not read; however, a single record containing a print code is read in place of the array control record. The print code determines the format for printing the values of the variable as defined by parameters. The print codes are the same as those used in an array control record. If any parameters of a given type are used, parameters must be used to define the corresponding variable for all layers in the model.

10. HK(NCOL,NROW) If there are any HK parameters, read only a print code.
11. [HANI(NCOL,NROW)] Include item 11 only if CHANI is less than or equal to 0. If there are any HANI parameters, read only a print code.
12. VKA(NCOL,NROW) If there are any VK or VANI parameters, read only a print code.
13. [Ss(NCOL,NROW)] Include item 13 only if at least one stress period is transient. If there are any SS parameters, read only a print code.
14. [Sy(NCOL,NROW)] Include item 14 only if at least one stress period is transient and LAYTYP is not 0. If there are any SY parameters, read only a print code.

15. [VKCB(NCOL,NROW)] Include item 15 only if LAYCBD (in the Discretization File) is not 0. If there are any VKCB parameters, read only a print code.
16. [WETDRY(NCOL,NROW)] Include item 16 only if LAYWET is not 0 and LAYTYP is not 0.

Explanation of Variables Read by the LPF Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

ILPFCB—is a flag and a unit number.

If ILPFCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control. The terms that are saved are storage, constant-head flow, and flow between adjacent cells.

If ILPFCB = 0, cell-by-cell flow terms will not be written.

If ILPFCB < 0, cell-by-cell flow for constant-head cells will be written in the listing file when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control. Cell-by-cell flow to storage and between adjacent cells will not be written to any file.

HDRY—is the head that is assigned to cells that are converted to dry during a simulation. Although this value plays no role in the model calculations, it is useful as an indicator when looking at the resulting heads that are output from the model. HDRY is thus similar to HNOFLO in the Basic Package, which is the value assigned to cells that are no-flow cells at the start of a model simulation.

NPLPF—is the number of LPF parameters.

LAYTYP—contains a flag for each layer that specifies the layer type. Use as many records as needed to enter a value for each layer.

0—confined

not 0—convertible.

LAYAVG—contains a flag for each layer that defines the method of calculating interblock transmissivity. Use as many records as needed to enter a value for each layer.

0—harmonic mean

1—logarithmic mean

2—arithmetic mean of saturated thickness and logarithmic-mean hydraulic conductivity.

CHANI—contains a value for each layer that is a flag or the horizontal anisotropy. If CHANI is less than or equal to 0, then variable HANI defines horizontal anisotropy. If CHANI is greater than 0, then CHANI is the horizontal anisotropy for the entire layer, and HANI is not read. If any HANI parameters are used, CHANI for all layers must be less than or equal to 0. Use as many records as needed to enter a value of CHANI for each layer.

LAYVKA—contains a flag for each layer that indicates whether variable VKA is vertical hydraulic conductivity or the ratio of horizontal to vertical hydraulic conductivity. Use as many records as needed to enter a value for each layer.

0—indicates VKA is vertical hydraulic conductivity

not 0—indicates VKA is the ratio of horizontal to vertical hydraulic conductivity, where the horizontal hydraulic conductivity is specified as HK in item 10.

LAYWET—contains a flag for each layer that indicates if wetting is active. Use as many records as needed to enter a value for each layer.

0—indicates wetting is inactive
not 0—indicates wetting is active

WETFCT—is a factor that is included in the calculation of the head that is initially established at a cell when it is converted from dry to wet. (See IHDWET.)

IWETIT—is the iteration interval for attempting to wet cells. Wetting is attempted every IWETIT iteration. If using the PCG solver (Hill, 1990), this applies to outer iterations, not inner iterations. If $IWETIT \leq 0$, it is changed to 1.

IHDWET—is a flag that determines which equation is used to define the initial head at cells that become wet:

If IHDWET = 0, equation 33A is used:
$$h = BOT + WETFCT (h_n - BOT) .$$

If IHDWET is not 0, equation 33B is used:
$$h = BOT + THRESH(WETDRY) .$$

PARNAM—is the name of a parameter to be defined. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter to be defined. For the LPF Package, the allowed parameter types are:

HK—defines variable HK, horizontal hydraulic conductivity
HANI—defines variable HANI, horizontal anisotropy
VK—defines variable VKA for layers for which VKA represents vertical hydraulic conductivity
(LAYVKA=0)
VANI—defines variable VKA for layers for which VKA represents vertical anisotropy (LAYVKA≠0)
SS—defines variable Ss, the specific storage
SY—defines variable Sy, the specific yield
VKCB—defines variable VKCB, the vertical hydraulic conductivity of a Quasi-three-dimensional confining layer.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NCLU—is the number of clusters required to define the parameter. Each repetition of Item 9 is a cluster (variables Layer, Mltarr, Zonarr, and IZ). There is usually only one cluster for each layer that is associated with a parameter. For example, if the parameter applies to cells in a single layer, there will generally be just one cluster. However, it is acceptable to have more than one cluster for the same layer.

Layer—is the layer number to which a cluster definition applies.

Mltarr—is the name of the multiplier array to be used to define variable values that are associated with a parameter. The name “NONE” means that there is no multiplier array, and the variable values will be set equal to Parval.

Zonarr—is the name of the zone array to be used to define the cells that are associated with a parameter. The name “ALL” means that there is no zone array, and all cells in the specified layer are part of the parameter.

IZ—is up to 10 zone numbers (separated by spaces) that define the cells that are associated with a parameter. These values are not used if ZONARR is specified as “ALL”. Values can be positive or negative, but 0 is not allowed. The end of the line, a zero value, or a non-numeric entry terminates the list of values.

HK—is the hydraulic conductivity along rows. HK is multiplied by horizontal anisotropy (see CHANI and HANI) to obtain hydraulic conductivity along columns.

HANI—is the ratio of hydraulic conductivity along columns to hydraulic conductivity along rows, where HK of item 10 specifies the hydraulic conductivity along rows. Thus, the hydraulic conductivity along columns is the product of the values in HK and HANI. Read only if CHANI \leq 0.

VKA—is either vertical hydraulic conductivity or the ratio of horizontal to vertical hydraulic conductivity depending on the value of LAYVKA. If LAYVKA is 0, VKA is vertical hydraulic conductivity. If LAYVKA is not 0, VKA is the ratio of horizontal to vertical hydraulic conductivity. In this case, HK is divided by VKA to obtain vertical hydraulic conductivity, and values of VKA typically are greater than or equal to 1.0.

Ss—is specific storage. Read only for a transient simulation (at least one transient stress period).

Sy—is specific yield. Read only for a transient simulation (at least one transient stress period) and if the layer is convertible (LAYTYP is not 0).

VKCB—is the vertical hydraulic conductivity of a Quasi-three-dimensional confining bed below a layer. Read only if there is a confining bed. Because there cannot be a confining bed beneath the bottom layer, VKCB cannot be specified for the bottom layer.

WETDRY—is a combination of the wetting threshold and a flag to indicate which neighboring cells can cause a cell to become wet. If WETDRY $<$ 0, only the cell below a dry cell can cause the cell to become wet. If WETDRY $>$ 0, the cell below a dry cell and the four horizontally adjacent cells can cause a cell to become wet. If WETDRY is 0, the cell cannot be wetted. The absolute value of WETDRY is the wetting threshold. When the sum of BOT and the absolute value of WETDRY at a dry cell is equaled or exceeded by the head at an adjacent cell, the cell is wetted. Read only if LAYTYP is not 0 and LAYWET is not 0.

Horizontal Flow Barrier Package

The Horizontal Flow Barrier Package is documented in Hsieh and Freckleton (1993). Input for the Horizontal Flow Barrier (HFB) Package is read from the file that has file type “HFB6” in the name file. The file type has been changed from that used in MODFLOW-96, which was “HFB”, because the input data are not compatible. All variables are read in free format.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—“#” must be in column 1. Item 0 can be repeated multiple times.

1. NPHFB MXFB NHFBNP

2. PARNAM PARTY Parval NLST

3. Layer IROW1 ICOL1 IROW2 ICOL2 Factor

NLST repetitions of Item 3 are required; they are read by module ULSTRD. (SFAC of the ULSTRD utility module applies to Factor).

Repeat Items 2 and 3 for each NPHFB parameter. Items 2 and 3 are not read if NPHFB is negative or zero.

FOR EACH STRESS PERIOD

4. Layer IROW1 ICOL1 IROW2 ICOL2 Hydchr

NHFBNP repetitions of Item 4 are read. Item 4 is not read if NHFBNP is negative or zero.

5. NACTHFB

6. Pname

NACTHFB repetitions of Item 6 are read. Item 6 is not read if NACTHFB is negative or zero.

Explanation of Variables Read by the HFB Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPHFB—is the number of horizontal-flow barrier parameters to be defined in Items 2 and 3. Note that for an HFB parameter to have an effect in the simulation, it must be defined in Items 2 and 3 and made active using Item 6.

MXFB—is the maximum number of HFB barriers that will be defined using parameters.

NHFBNP—is the number of HFB barriers not defined by parameters.

PARNAM—is the name of a parameter. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter. For the HFB Package, the only allowed parameter type is HFB, which defines values of the hydraulic characteristic of the barrier.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NLST—is the number of horizontal-flow barrier cells included in the parameter.

Layer—is the number of the model layer in which the horizontal-flow barrier is located.

IROW1—is the row number of the cell on one side of the horizontal-flow barrier.

ICOL1—is the column number of the cell on one side of the horizontal-flow barrier.

IROW2—is the row number of the cell on the other side of the horizontal-flow barrier.

ICOL2—is the column number of the cell on the other side of the horizontal-flow barrier.

Factor—is the factor used to calculate hydraulic characteristic from the parameter value. The hydraulic characteristic is the product of Factor and the parameter value.

Hydchr—is the hydraulic characteristic of the horizontal-flow barrier. The hydraulic characteristic is the barrier hydraulic conductivity divided by the width of the horizontal-flow barrier.

NACTHFB—is the number of active HFB parameters.

Pname—is the name of a parameter to be used in the simulation. NACTHFB parameter names will be read.

River Package

Input to the River (RIV) Package is read from the file that has file type "RIV" in the name file. Optional variables are shown in brackets. All variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the non-optional variables have 10-character fields and the optional variables are free format.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. [**PARAMETER** NPRIV MXL]

This optional item must start with the word "PARAMETER".

2. MXACTR IRIVCB [Option]

3. [PARNAM PARTYP Parval NLST]

4. Layer Row Column Stage Condfact Rbot [xyz]

NLST repetitions of Item 4 are required; they are read by module ULSTRD. (SFAC of the ULSTRD utility module applies to Condfact).

Repeat Items 3 and 4 for each NPRIV parameter.

FOR EACH STRESS PERIOD

5. ITMP NP

6. Layer Row Column Stage Cond Rbot [xyz]

ITMP repetitions of Item 6 are read by module ULSTRD if ITMP > 0. (SFAC of the ULSTRD utility module applies to Cond.) Item 6 is not read if ITMP is negative or 0.

7. [Pname]

(Item 7 is repeated NP times. It is not read if NP is negative or 0.)

Explanation of Variables Read by the RIV Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPRIV—is the number of river parameters.

MXL—is the maximum number of river reaches that will be defined using parameters.

MXACTR—is the maximum number of river reaches in use during any stress period, including those that are defined using parameters.

IRIVCB—is a flag and a unit number.

If IRIVCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

If IRIVCB = 0, cell-by-cell flow terms will not be written.

If IRIVCB < 0, river leakage for each reach will be written to the listing file when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

Option—is an optional list of character values.

“AUXILIARY abc” or “AUX abc”—defines an auxiliary variable, named "abc", which will be read for each river reach as part of Items 4 and 6. Up to five variables can be specified, each of which must be preceded by "AUXILIARY" or "AUX." These variables will not be used by the Ground-Water Flow Process Package, but they will be available for use by other processes. The auxiliary variable values will be read after the Rbot variable.

“CBCALLOCATE” or “CBC”—indicates that memory should be allocated to store cell-by-cell flow for each river reach in order to make these flows available for use in other packages.

PARNAM—is the name of a parameter. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter. For the RIV Package, the only allowed parameter type is RIV, which defines values of riverbed hydraulic conductance.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NLST—is the number of river reaches that are included in the parameter.

ITMP—is a flag and a counter.

If $ITMP < 0$, non-parameter river data from the last stress period will be reused.

If $ITMP \geq 0$, ITMP will be the number of non-parameter reaches read for the current stress period.

NP—is the number of parameters in use in the current stress period.

Layer—is the layer number of the cell containing the river reach.

Row—is the row number of the cell containing the river reach.

Column—is the column number of the cell containing the river reach.

Stage—is the head in the river.

Condfact—is the factor used to calculate riverbed hydraulic conductance from the parameter value. The conductance is the product of Condfact and the parameter value.

Cond—is the riverbed hydraulic conductance.

Rbot—is the elevation of the bottom of the riverbed.

[xyz]—represents any auxiliary variables for a river reach that have been defined in Item 2. The auxiliary variables must be present in each repetition of Items 4 and 6 if they are defined in Item 2.

Pname—is the name of a parameter that is being used in the current stress period. NP parameter names will be read.

Recharge Package

Input to the Recharge (RCH) Package is read from the file that has type "RCH" in the name file. All single-valued variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the variables have 10-character fields.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. [**PARAMETER** NPRCH]

This optional item must start with the word "PARAMETER".

2. NRCHOP IRCHCB

3. [PARNAM PARTYP Parval NCLU]

4. [Mltarr Zonarr IZ]

Repeat Item 4 NCLU times. Each repetition of Item 4 is called a parameter cluster.

Repeat Items 3 and 4 for each parameter to be defined (that is, NPRCH times).

FOR EACH STRESS PERIOD

5. INRECH INIRCH

6. [RECH(NCOL,NROW)] -- U2DREL if NPRCH=0 and if INRECH ≥ 0

7. [Pname [IRCHPF]] -- if NPRCH > 0 and if INRECH > 0

Either Item 6 or Item 7 may be read, but not both. If Item 7 is read, it is repeated INRECH times.

8. [IRCH(NCOL,NROW)] -- U2DINT If NRCHOP=2 and if INIRCH ≥ 0

Explanation of Variables Read by the RCH Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPRCH—is the number of recharge parameters.

NRCHOP—is the recharge option code. Recharge fluxes are defined in a layer variable, RECH, with one value for each vertical column. Accordingly, recharge is applied to one cell in each vertical column, and the option code determines which cell in the column is selected for recharge.

1—Recharge is only to the top grid layer.

2—Vertical distribution of recharge is specified in layer variable IRCH.

3—Recharge is applied to the highest active cell in each vertical column. A constant-head node intercepts recharge and prevents deeper infiltration.

IRCHCB—is a flag and a unit number.

If IRCHCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

If IRCHCB ≤ 0, cell-by-cell flow terms will not be written.

PARNAM—is the name of a parameter to be defined. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter to be defined. For the RCH Package, the only allowed parameter type is RCH, which defines values of the recharge flux.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NCLU—is the number of clusters required to define a parameter. Each repetition of Item 4 is a cluster (variables Mltarr, Zonarr, and IZ). There is usually only one cluster used to define a RCH parameter; however, it is acceptable to have more than one cluster.

Mltarr—is the name of the multiplier array to be used to define cell values that are determined by a parameter. The name “NONE” means that there is no multiplier array, and the cell values will be set equal to Parval.

Zonarr—is the name of the zone array to be used to define the cells that are associated with a parameter. The name “ALL” means that there is no zone array, and all cells in the layer are associated with the parameter.

IZ—is up to 10 zone numbers (separated by spaces) that define the cells that are associated with a parameter. These values are not used if Zonarr is specified as “ALL.” Values can be positive or negative, but 0 is not allowed. The end of the line, a zero value, or a non-numeric entry terminates the list of values.

INRECH—is the RECH read flag. Its function depends on whether or not parameters are being used.

If no parameters are being used ($\text{NPRCH} = 0$):

If $\text{INRECH} \geq 0$, a layer variable of recharge fluxes, RECH, is read.

If $\text{INRECH} < 0$, recharge rates from the preceding stress period are used.

If parameters are being used ($\text{NPRCH} > 0$):

If $\text{INRECH} > 0$, INRECH is the number of parameters that will be used to define RECH in the current stress period. Item 7 defines the names of the parameters.

If $\text{INRECH} < 0$, recharge parameters from the preceding stress period are used.

$\text{INRECH} = 0$ is not allowed. That is, when parameters are used, at least one parameter must be specified each stress period.

INIRCH—is the IRCH read flag, which is read only if NRCHOP is two:

If $\text{INIRCH} \geq 0$, a layer variable of layer numbers (IRCH) is read.

If $\text{INIRCH} < 0$, the variable (IRCH) used in the preceding stress period is reused.

RECH—is the recharge flux (LT^{-1}). Read only if INRECH is greater than or equal to zero and if $\text{NPRCH} = 0$.

Pname—is the name of a parameter that will be used to define the RECH variable in the current stress period. Read INRECH values if $\text{NPRCH} > 0$ and $\text{INRECH} > 0$.

IRCHPF—is an optional format code for printing the RECH variable after it has been defined by parameters. The format codes are the same as those used in the U2DREL array reading utility module.

IRCH—is the layer number variable that defines the layer in each vertical column where recharge is applied. Read only if NRCHOP is two and if INIRCH is greater than or equal to zero.

Well Package

Input to the Well (WEL) Package is read from the file that has type "WEL" in the name file. Optional variables are shown in brackets. All variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the non-optional variables have 10-character fields and the optional variables are free format.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. [**PARAMETER** NPWEL MXL]

This optional item must start with the word "PARAMETER".

2. MXACTW IWELCB [Option]

3. [PARNAM PARTYP Parval NLST]

4. Layer Row Column Qfact [xyz]

NLST repetitions of Item 4 are required; they are read by module ULSTRD. (SFAC of the ULSTRD utility module applies to Qfact). Repeat Items 3 and 4 for each NPWEL parameter.

FOR EACH STRESS PERIOD

5. ITMP NP

6. Layer Row Column Q [xyz]

(ITMP repetitions of Item 6 are read by module ULSTRD if ITMP > 0. (SFAC of the ULSTRD utility module applies to Q.) Item 6 is not read if ITMP is negative or zero.

7. [Pname]

(Item 7 is repeated NP times. It is not read if NP is negative or 0.)

Explanation of Variables Read by the WEL Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPWEL—is the number of well parameters.

MXL—is the maximum number of wells that will be defined using parameters.

MXACTW—is the maximum number of wells in use during any stress period, including those that are defined using parameters.

IWELCB—is a flag and a unit number.

If IWELCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

If IWELCB = 0, cell-by-cell flow terms will not be written.

If IWELCB < 0, well recharge for each well will be written to the listing file when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

Option—is an optional list of character values.

“AUXILIARY abc” or “AUX abc”—defines an auxiliary variable, named "abc", which will be read for each well as part of Items 4 and 6. Up to five variables can be specified, each of which must be preceded by "AUXILIARY" or "AUX." These variables will not be used by the Ground-Water Flow Process, but they will be available for use by other processes. The auxiliary variable values will be read after the Q variable.

“CBCALLOCATE” or “CBC”—indicates that memory should be allocated to store cell-by-cell flow for each well in order to make these flows available for use in other packages.

PARNAM—is the name of a parameter. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter. For the WEL Package, the only allowed parameter type is Q, which defines values of the volumetric recharge rate.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NLST—is the number of wells that are included in the parameter.

ITMP—is a flag and a counter.

If $ITMP < 0$, non-parameter well data from the last stress period will be reused.

If $ITMP \geq 0$, ITMP will be the number of non-parameter wells read for the current stress period.

NP—is the number of parameters in use in the current stress period.

Layer—is the layer number of the model cell that contains the well.

Row—is the row number of the model cell that contains the well.

Column—is the column number of the model cell that contains the well.

Qfact—is the factor used to calculate well recharge rate from the parameter value. The recharge rate is the product of Qfact and the parameter value.

Q—is the volumetric recharge rate. A positive value indicates recharge and a negative value indicates discharge (pumping).

[xyz]—represents any auxiliary variables for a well that have been defined in Item 2. The auxiliary variables must be present in each repetition of Items 4 and 6 if they are defined in Item 2.

Pname—is the name of a parameter that is being used in the current stress period. NP parameter names will be read.

Drain Package

Input to the Drain (DRN) Package is read from the file that has type "DRN" in the name file. Optional variables are shown in brackets. All variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the non-optional variables have 10-character fields and the optional variables are free format.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. [**PARAMETER** NPDRN MXL]

This optional item must start with the word "PARAMETER".

2. MXACTD IDRNCB [Option]

3. [PARNAM PARTYP Parval NLST]

4. Layer Row Column Elevation Condfact [xyz]

NLST repetitions of Item 4 are required; they are read by module ULSTRD. (SFAC of the ULSTRD utility module applies to Condfact).

Repeat Items 3 and 4 for each NPDRN parameter.

FOR EACH STRESS PERIOD

5. ITMP NP

6. Layer Row Column Elevation Cond [xyz]

ITMP repetitions of Item 6 are read by module ULSTRD if ITMP > 0. (SFAC of the ULSTRD utility module applies to Cond.) Item 6 is not read if ITMP is negative or 0.

7. [Pname]

(Item 7 is repeated NP times. It is not read if NP is negative or 0.)

Explanation of Variables Read by the DRN Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPDRN—is the number of drain parameters.

MXL—is the maximum number of drain cells that will be defined using parameters.

MXACTD—is the maximum number of drain cells in use during any stress period, including those that are defined using parameters.

IDRNCB—is a flag and a unit number.

If IDRNCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

If IDRNCB = 0, cell-by-cell flow terms will not be written.

If IDRNCB < 0, drain leakage for each drain cell will be written to the listing file when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

Option—is an optional list of character values.

“AUXILIARY abc” or “AUX abc”—defines an auxiliary variable, named "abc", which will be read for each drain as part of Items 4 and 6. Up to five variables can be specified, each of which must be preceded by "AUXILIARY" or "AUX." These variables will not be used by the Ground-Water Flow Process, but they will be available for use by other processes. The auxiliary variable values will be read after the Cond variable.

CBCALLOCATE or CBC—indicates that memory should be allocated to store cell-by-cell flow for each drain in order to make these flows available for use in other packages.

PARNAM—is the name of a parameter. This name can consist of 1 to 10 characters and is not case sensitive. That is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter. For the DRN Package, the only allowed parameter type is DRN, which defines values of the drain hydraulic conductance.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NLST—is the number of drain cells that are included in the parameter.

ITMP—is a flag and a counter.

If $ITMP < 0$, non-parameter drain data from the last stress period will be reused.

If $ITMP \geq 0$, ITMP will be the number of non-parameter drains read for the current stress period.

NP—is the number of parameters in use in the current stress period.

Layer—is the layer number of the cell containing the drain.

Row—is the row number of the cell containing the drain.

Column—is the column number of the cell containing the drain.

Elevation—is the elevation of the drain.

Condfact—is the factor used to calculate drain hydraulic conductance from the parameter value. The conductance is the product of Condfact and the parameter value.

Cond—is the hydraulic conductance of the interface between the aquifer and the drain.

[xyz]—represents any auxiliary variables for a drain that have been defined in Item 2. The auxiliary variables must be present in each repetition of Items 4 and 6 if they are defined in Item 2.

Pname—is the name of a parameter that is being used in the current stress period. NP parameter names will be read.

Evapotranspiration Package

Input to the Evapotranspiration (EVT) Package is read from the file that is type "EVT" in the name file. All single-valued variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the variables have 10-character fields.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. [**PARAMETER** NPEVT]
This optional item must start with the word "PARAMETER".
2. NEVTOP IEVTCB
3. [PARNAM PARTYP Parval NCLU]
4. [Mltarr Zonarr IZ]
Repeat Item 4 NCLU times. Each repetition of Item 4 is called a parameter cluster.
Repeat Items 3-4 for each parameter to be defined (that is, NPEVT times).

FOR EACH STRESS PERIOD

5. INSURF INEVTR INEXDP INIEVT
6. [SURF(NCOL,NROW)] -- U2DREL If INSURF \geq 0
7. [EVTR(NCOL,NROW)] -- U2DREL If NPEVT = 0 and if INEVTR \geq 0
8. [Pname [IEVTPF]] -- if NPEVT > 0 and if INEVTR > 0
Either Item 7 or Item 8 may be read, but not both. If Item 8 is read, it is repeated INEVTR times.
9. [EXDP(NCOL,NROW)] -- U2DREL If INEXDP \geq 0
10. [IEVT(NCOL,NROW)] -- U2DINT If NEVTOP = 2 and if INIEVT \geq 0

Explanation of Variables Read by the EVT Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPEVT—is the number of evapotranspiration parameters.

NEVTOP—is the evapotranspiration (ET) option code. ET variables (ET surface, maximum ET rate, and extinction depth) are specified in layer variables, SURF, EVTR, and EXDP, with one value for each vertical column. Accordingly, ET is calculated for one cell in each vertical column. The option codes determine the cell within a column for which ET will be calculated.

1—ET is calculated only for cells in the top grid layer.

2—The cell for each vertical column is specified by the user in variable IEVT.

IEVTCB—is a flag and a unit number.

If IEVTCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

If IEVTCB \leq 0, cell-by-cell flow terms will not be written.

PARNAM—is the name of a parameter to be defined. This name can consist of 1 to 10 characters and is not case sensitive; that is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter to be defined. For the EVT Package, the only allowed parameter type is EVT, which defines values of the maximum ET flux.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NCLU—is the number of clusters required to define a parameter. Each repetition of Item 4 is a cluster (variables Mltarr, Zonarr, and IZ). There is usually only one cluster used to define an EVT parameter; however, it is acceptable to have more than one cluster.

Mltarr—is the name of the multiplier array to be used to define the values that are determined by a parameter. The name “NONE” means that there is no multiplier array, and the values will be set equal to Parval.

Zonarr—is the name of the zone array to be used to define the cells that are associated with a parameter. The name “ALL” means that there is no zone array, and all cells are associated with the parameter.

IZ—is up to 10 zone numbers (separated by spaces) that define the cells that are associated with a parameter. These values are not used if Zonarr is specified as “ALL.” Values can be positive or negative, but 0 is not allowed. The end of the line, a zero value, or a non-numeric entry terminates the list of values.

INSURF—is the ET surface (SURF) read flag.

If $INSURF \geq 0$, a layer variable containing the ET surface elevation (SURF) will be read.

If $INSURF < 0$, the ET surface from the preceding stress period will be reused.

INEVTR—is the EVTR read flag. Its function depends on whether or not parameters are being used.

If no parameters are being used ($NPEVT=0$):

If $INEVTR \geq 0$, a layer variable containing the maximum ET rate (EVTR) will be read.

If $INEVTR < 0$, the maximum ET rate from the preceding stress period will be reused.

If parameters are being used ($NPEVT>0$):

If $INEVTR > 0$, INEVTR is the number of parameters that will be used to define EVTR in the current stress period. Item 8 defines the names of the parameters.

If $INEVTR < 0$, EVT parameters from the preceding stress period are used.

$INEVTR = 0$ is not allowed. That is, when parameters are used, at least one parameter must be specified each stress period

INEXDP—is the extinction depth (EXDP) read flag.

If $INEXDP \geq 0$, a layer variable containing the extinction depth (EXDP) will be read.

If $INEXDP < 0$, the extinction depth from the preceding stress period will be reused.

INIEVT—is the layer indicator (IEVT) read flag. It is read only if the ET option (NEVTOP) is equal to two.

If $INIEVT \geq 0$, a layer variable containing the layer indicators (IEVT) will be read.

If $INIEVT < 0$, layer indicators used during the preceding stress period will be reused.

SURF—is the elevation of the ET surface. This variable is read only if $INSURF \geq 0$

EVTR—is the maximum ET flux [volumetric flow rate per unit area (LT^{-1})]. This variable is read only if $INEVTR \geq 0$ and if $NPEVT=0$.

Pname—is the name of a parameter that will be used to define the EVTR variable in the current stress period. Read INEVTR values if $NPEVT > 0$ and $INEVTR > 0$.

IEVTPF—is an optional format code for printing the EVTR variable after it has been defined by parameters. The format codes are the same as those used in the U2DREL array reading utility module.

EXDP—is the ET extinction depth. This variable is read only if $INEXDP \geq 0$.

IEVT—is the layer indicator variable. For each horizontal location, it indicates the layer from which ET is removed. It is read only if the ET option is equal to two and if $INIEVT \geq 0$.

General-Head Boundary Package

Input to the General-Head Boundary (GHB) Package is read from the file that has file type "GHB" in the name file. Optional variables are shown in brackets. All variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the non-optional variables have 10-character fields and the optional variables are free format.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. [**PARAMETER** NPGHB MXL]
This optional item must start with the word "PARAMETER".
2. MXACTB IGHBCB [Option]
3. [PARNAM PARTYP Parval NLST]
4. Layer Row Column Bhead Condfact [xyz]
NLST repetitions of Item 4 are required; they are read by module ULSTRD. (SFAC of the ULSTRD utility module applies to Condfact).
Repeat Items 3 and 4 for each NPGHB parameter.

FOR EACH STRESS PERIOD

5. ITMP NP
6. Layer Row Column Bhead Cond [xyz]
ITMP repetitions of Item 6 are read by module ULSTRD if ITMP > 0. (SFAC of the ULSTRD utility module applies to Cond.) Item 6 is not read if ITMP is negative or 0.
7. [Pname]
(Item 7 is repeated NP times. It is not read if NP is negative or 0.)

Explanation of Variables Read by the GHB Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPGHB—is the number of general-head boundary parameters.

MXL—is the maximum number of general-head-boundary cells that will be defined using parameters.

MXACTB—is the maximum number of general-head boundary cells in use during any stress period, including those that are defined using parameters.

IGHBCB—is a flag and a unit number.

If IGHBCB > 0, it is the unit number to which cell-by-cell flow terms will be written when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

If IGHBCB = 0, cell-by-cell flow terms will not be written.

If IGHBCB < 0, boundary leakage for each GHB cell will be written to the listing file when "SAVE BUDGET" or a non-zero value for ICBCFL is specified in Output Control.

Option—is an optional list of character values.

“AUXILIARY abc” or “AUX abc”—defines an auxiliary variable, named "abc", which will be read for each general-head boundary as part of Items 4 and 6. Up to five variables can be specified, each of which must be preceded by "AUXILIARY" or "AUX." These variables will not be used by the Ground-Water Flow Process, but they will be available for use by other processes. The auxiliary variable values will be read after the Cond variable.

“CBCALLOCATE” or “CBC”—indicates that memory should be allocated to store cell-by-cell flow for each general-head boundary in order to make these flows available for use in other packages.

PARNAM—is the name of a parameter. This name can consist of 1 to 10 characters and is not case sensitive; that is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter to be defined. For the GHB Package, the only allowed parameter type is GHB, which defines values of the general-head boundary hydraulic conductance.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NLST—is the number of head-dependent boundary cells that are included in the parameter.

ITMP—is a flag and a counter.

If $ITMP < 0$, non-parameter GHB data from the preceding stress period will be reused.

If $ITMP \geq 0$, ITMP is the number of non-parameter general-head boundaries read for the current stress period.

NP—is the number of parameters in use in the current stress period.

Layer—is the layer number of the cell affected by the head-dependent boundary.

Row—is the row number of the cell affected by the head-dependent boundary.

Column—is the column number of the cell affected by the head-dependent boundary.

Bhead—is the head on the boundary.

Condfact—is the factor used to calculate hydraulic conductance from the parameter value. The conductance is the product of Condfact and the parameter value.

Cond—is the hydraulic conductance of the interface between the aquifer cell and the boundary.

[xyz]—represents any auxiliary variables for a boundary that have been defined in Item 2. The auxiliary variables must be present in each repetition of Items 4 and 6 if they are defined in item 2.

Pname—is the name of a parameter that is being used in the current stress period. NP parameter names will be read.

Constant-Head Boundary Package

Input to the Constant-Head Boundary (CHD) Package is read from the file that has file type "CHD" in the name file. Optional variables are shown in brackets. All variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, the non-optional variables have 10-character fields and the optional variables are free format.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. [**PARAMETER** NPCHD MXL]
This optional item must start with the word "PARAMETER".
2. MXACTC [Option]
3. [PARNAM PARTYP Parval NLST]
4. Layer Row Column Shdfact Eheadfact [xyz]
NLST repetitions of Item 4 are required; they are read by module ULSTRD. (SFAC of the ULSTRD utility module applies to Shdfact and Eheadfact).
Repeat Items 3 and 4 for each NPGHB parameter.

FOR EACH STRESS PERIOD

5. ITMP NP
6. Layer Row Column Shead Ehead [xyz]
ITMP repetitions of Item 6 are read by module ULSTRD if ITMP > 0. (SFAC of the ULSTRD utility module applies to Shead and Ehead.) Item 6 is not read if ITMP is negative or 0.
7. [Pname]
(Item 7 is repeated NP times. It is not read if NP is negative or 0.)

Explanation of Variables Read by the CHD Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

NPCHD—is the number of constant-head boundary parameters.

MXL—is the maximum number of constant-head-boundary cells that will be defined using parameters.

MXACTC—is the maximum number of constant-head boundary cells in use during any stress period, including those that are defined using parameters.

Option—is an optional list of character values.

"AUXILIARY abc" or "AUX abc"—defines an auxiliary variable, named "abc", which will be read for each constant-head boundary as part of Items 4 and 6. Up to five variables can be specified, each of which must be preceded by "AUXILIARY" or "AUX." These variables will not be used by the Ground-Water Flow Process, but they will be available for use by other processes. The auxiliary variable values will be read after the Ehead variable.

PARNAM—is the name of a parameter. This name can consist of 1 to 10 characters and is not case sensitive; that is, any combination of the same characters with different case will be equivalent.

PARTYP—is the type of parameter to be defined. For the CHD Package, the only allowed parameter type is CHD, which defines values of the start and end head at the boundary.

Parval—is the parameter value. This parameter value may be overridden by a value in the Sensitivity Process input file or by a value generated by the Parameter-Estimation Process.

NLST—is the number of constant-head cells that are included in the parameter.

ITMP—is a flag and a counter.

If $ITMP < 0$, non-parameter CHD data from the preceding stress period will be reused.

If $ITMP \geq 0$, ITMP is the number of non-parameter constant-head boundaries read for the current stress period.

NP—is the number of parameters in use in the current stress period.

Layer—is the layer number of the constant-head boundary.

Row—is the row number of the constant-head boundary.

Column—is the column number of the constant-head boundary.

Shdfact—is the factor used to calculate the head at the boundary at the start of the stress period from the parameter value. The head is the product of Shdfact and the parameter value.

Ehdfact—is the factor used to calculate the head at the boundary at the end of the stress period from the parameter value. The head is the product of Ehdfact and the parameter value.

Shead—is the head at the boundary at the start of the stress period.

Ehead—is the head at the boundary at the end of the stress period.

[xyz]—represents any auxiliary variables for a constant-head boundary that have been defined in Item 2. The auxiliary variables must be present in each repetition of Items 4 and 6 if they are defined in Item 2.

Pname—is the name of a parameter that is being used in the current stress period. NP parameter names will be read.

Strongly Implicit Procedure Package

Input to the Strongly Implicit Procedure (SIP) Package is read from the file that is type "SIP" in the name file. All numeric variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, all the variables have 10-character fields.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. MXITER NPARM
2. ACCL HCLOSE IPCALC WSEED IPRSIP

Explanation of Variables Read by the SIP Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

MXITER—is the maximum number of times through the iteration loop in one time step in an attempt to solve the system of finite-difference equations.

NPARM—is the number of iteration variables to be used. Five variables are generally sufficient.

ACCL—is the acceleration variable. It must be greater than zero and is generally equal to one. If a zero is entered, it is changed to one.

HCLOSE—is the head change criterion for convergence. When the maximum absolute value of head change from all nodes during an iteration is less than or equal to HCLOSE, iteration stops.

IPCALC—is a flag indicating where the seed for calculating iteration variables will come from.

0—the seed entered by the user will be used.

1—the seed will be calculated at the start of the simulation from problem variables.

WSEED—is the seed for calculating iteration variables. It is always read, but it is used only if IPCALC is equal to zero.

IPRSIP—is the printout interval for SIP. If IPRSIP is equal to zero, it is changed to 999. The maximum head change (positive or negative) is printed for each iteration of a time step whenever the time step is an even multiple of IPRSIP. This printout also occurs at the end of each stress period regardless of the value of IPRSIP.

Slice-Successive Overrelaxation Package

Input to the Slice-Successive Overrelaxation (SOR) Package is read from the file that is type "SOR" in the name file. All numeric variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, all the variables have 10-character fields.

FOR EACH SIMULATION

0. [#Text]

Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.

1. MXITER

2. ACCL HCLOSE IPRSOR

Explanation of Variables Read by the SOR Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

MXITER—is the maximum number of iterations allowed in a time step.

ACCL—is the acceleration variable, usually between 1.0 and 2.0.

HCLOSE—is the head change criterion for convergence. When the maximum absolute value of head change from all nodes during an iteration is less than or equal to HCLOSE, iteration stops.

IPRSOR—is the printout interval for SOR. IF IPRSOR is equal to zero, it is changed to 999. The maximum head change (positive or negative) is printed for each iteration of a time step whenever the time step is an even multiple of IPRSOR. This printout also occurs at the end of each stress period regardless of the value of IPRSOR.

Preconditioned Conjugate-Gradient Package

Input to the Preconditioned Conjugate-Gradient (PCG) Package is read from the file that is type "PCG" in the name file. All numeric variables are free format if the option "FREE" is specified in the Basic Package input file; otherwise, all the variables have 10-character fields.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. MXITER ITER1 NPCOND
2. HCLOSE RCLOSE RELAX NPBOL IPRPCG MUTPCG DAMP

Explanation of Variables Read by the PCG Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The "#" character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

MXITER—is the maximum number of outer iterations—that is, calls to the solution routine. For a linear problem MXITER should be 1, unless more than 50 inner iterations are required, when MXITER could be as large as 10. A larger number (generally less than 100) is required for a nonlinear problem.

ITER1—is the number of inner iterations. For nonlinear problems, ITER1 usually ranges from 10 to 30; a value of 30 will be sufficient for most linear problems.

NPCOND—is the flag used to select the matrix conditioning method:

- 1—is for Modified Incomplete Cholesky (for use on scalar computers)
- 2—is for Polynomial (for use on vector computers or to conserve computer memory)

HCLOSE—is the head change criterion for convergence, in units of length. When the maximum absolute value of head change from all nodes during an iteration is less than or equal to HCLOSE, and the criterion for RCLOSE is also satisfied (see below), iteration stops.

RCLOSE—is the residual criterion for convergence, in units of cubic length per time. When the maximum absolute value of the residual at all nodes during an iteration is less than or equal to RCLOSE, and the criterion for HCLOSE is also satisfied (see above), iteration stops.

For nonlinear problems, convergence is achieved when the convergence criteria are satisfied for the first inner iteration.

RELAX—is the relaxation parameter used with NPCOND = 1. Usually, RELAX = 1.0, but for some problems a value of 0.99, 0.98, or 0.97 will reduce the number of iterations required for convergence. RELAX is not used if NPCOND is not 1.

NPBOL—is used when NPCOND = 2 to indicate whether the estimate of the upper bound on the maximum eigenvalue is 2.0, or whether the estimate will be calculated. NPBOL = 2 is used to specify the value is 2.0; for any other value of NPBOL, the estimate is calculated. Convergence is generally insensitive to this parameter. NPBOL is not used if NPBOL is not 2.

IPRPCG—is the printout interval for PCG. If IPRPCG is equal to zero, it is changed to 999. The maximum head change (positive or negative) and residual change are printed for each iteration of a time step whenever the time

step is an even multiple of IPRPCG. This printout also occurs at the end of each stress period regardless of the value of IPRPCG.

MUTPCG—is a flag that controls printing of convergence information from the solver:

- 0—is for printing tables of maximum head change and residual each iteration
- 1—is for printing only the total number of iterations
- 2—is for no printing
- 3—is for printing only if convergence fails

DAMP—is the damping factor. It is typically set equal to one, which indicates no damping. A value less than 1 and greater than 0 causes damping.

Direct Solver Package Input Instructions

Input to the Direct Solver (DE4) Package is read from the file that is type "DE4" in the name file. All numeric variables are free format.

FOR EACH SIMULATION

0. [#Text]
Item 0 is optional—"#" must be in column 1. Item 0 can be repeated multiple times.
1. ITMX MXUP MXLOW MXBW
2. IFREQ MUTD4 ACCL HCLOSE IPRD4

Explanation of Variables Read by the DE4 Package

Text—is a character variable (199 characters) that starts in column 2. Any characters can be included in Text. The “#” character must be in column 1. Except for the name file, lines beginning with # are restricted to these first lines of the file. Text is printed when the file is read.

ITMX—is the maximum number of iterations each time step. Specify ITMX = 1 if iteration is not desired. Ideally iteration would not be required for direct solution; however, it is necessary to iterate if the flow equation is nonlinear (see explanation for IFREQ = 3) or if computer precision limitations result in inaccurate calculations as indicated by a large water budget error. For a nonlinear flow equation, each iteration is equally time consuming because [A] is changed each iteration and Gaussian elimination is required after each change. This is called external iteration. For a linear equation, iteration is significantly faster because [A] is changed at most once per time step; thus, Gaussian elimination is required at most once per time step. This is called internal iteration.

MXUP—is the maximum number of equations in the upper part of the equations to be solved. This value impacts the amount of memory used by the DE4 Package. If specified as 0, the program will calculate MXUP as half the number of cells in the model, which is an upper limit. The actual number of equations in the upper part will be less than half the number of cells whenever there are no-flow and constant-head cells because flow equations are not formulated for these cells. The DE4 Package prints the actual number of equations in the upper part when it runs. The printed value can be used for MXUP in future runs in order to minimize memory usage.

MXLOW—is the maximum number of equations in the lower part of equations to be solved. This value impacts the amount of memory used by the DE4 Package. If specified as 0, the program will calculate MXLOW as half the number of cells in the model, which is an upper limit. The actual number of equations in the lower part will be less than half the number of cells whenever there are no-flow and constant-head cells because flow equations are not formulated for these cells. The DE4 Package prints the actual number of equations in the lower part when it runs. The printed value can be used for MXLOW in future runs in order to minimize memory usage.

MXBW—is the maximum band width plus 1 of the [AL] matrix. This value impacts the amount of memory used by the DE4 Package. If specified as 0, the program will calculate MXBW as the product of the two smallest grid dimensions plus 1, which is an upper limit. The DE4 Package prints the actual band width plus 1 when it runs. The printed value can be used for MXBW in future runs in order to minimize memory usage.

IFREQ—is a flag indicating the frequency at which coefficients in [A] change. This affects the efficiency of solution; significant work can be avoided if it is known that [A] remains constant all or part of the time.

IFREQ = 1 indicates that the flow equations are linear and that coefficients of simulated head for all stress terms are constant for all stress periods. To meet the linearity requirement, all model layers must be confined (which is specified in the Block-Centered Flow Package by setting LAYCON equal to 0 for all layers or in the Layer-Property Flow Package by setting LAYTYP equal to 0 for all layers), and there

must be no formulations that change based upon head (such as seepage from a river changing from head dependent flow to a constant flow when head drops below the bottom of the riverbed). Examples of coefficients of simulated head for stress terms are riverbed conductance, drain conductance, maximum evapotranspiration rate, evapotranspiration extinction depth, and general-head boundary conductance.

IFREQ = 2 indicates that the flow equations are linear, but coefficients of simulated head for some stress terms may change at the start of each stress period. (See IFREQ = 1 for information about linear equations.) Examples of coefficients of simulated head for stress terms are riverbed conductance, drain conductance, maximum evapotranspiration rate, evapotranspiration extinction depth, and general-head boundary conductance. For a simulation consisting of only one stress period, IFREQ = 2 has the same meaning as IFREQ = 1.

IFREQ = 3 indicates that a nonlinear flow equation is being solved, which means that some terms in [A] depend on simulated head. Examples of head-dependent terms in [A] are transmissivity for water-table layers, which is based on saturated thickness; flow terms for rivers, drains, and evapotranspiration if they convert between head dependent flow and constant flow; and the change in storage coefficient when a cell converts between confined and unconfined. When a nonlinear flow equation is being solved, external iteration (ITMX > 1) is normally required to accurately approximate the nonlinearities. Note that when nonlinearities caused by water-table calculations are part of a simulation, there are not necessarily any obvious signs in the output from a simulation that does not use external iteration to indicate that iteration is needed. In particular, the budget error may be acceptably small without iteration even though there is significant error in head because of nonlinearity. To understand this, consider the water-table correction for transmissivity. Each iteration a new transmissivity is calculated based on the previous head. Then the flow equations are solved, and a budget is computed using the new head with the same transmissivities. No budget discrepancy results because heads are correct for the transmissivity being used at this point; however, the new heads may mean that there should be a significant change in transmissivity. The new transmissivity will not be calculated unless there is another iteration. Therefore, when one or more layers is under water-table conditions, iteration should always be tried. The maximum change in head each iteration (printed by DE4 when IPRD4 = 1 and MUTD4 = 0) provides an indication of the impact of all nonlinearities.

MUTD4—is a flag that indicates the quantity of information that is printed when convergence information is printed for a time step.

MUTD4 = 0 indicates that the number of iterations in the time step and the maximum head change each iteration are printed.

MUTD4 = 1 indicates that only the number of iterations in the time step is printed.

MUTD4 = 2 indicates no information is printed.

ACCL—is a multiplier for the computed head change for each iteration. Normally this value is 1. A value greater than 1 may be useful for improving the rate of convergence when using external iteration to solve nonlinear problems (IFREQ = 3). ACCL should always be 1 for linear problems. When ITMX = 1, ACCL is changed to 1 regardless of the input value; however, a value must always be specified.

HCLOSE—is the head change closure criterion. If iterating (ITMX > 1), iteration stops when the absolute value of head change at every node is less than or equal to HCLOSE. HCLOSE is not used if not iterating, but a value must always be specified.

IPRD4—is the time step interval for printing out convergence information when iterating (ITMX > 1). For example, if IPRD4 is 2, convergence information is printed every other time step. A value must always be specified even if not iterating.

Input Instructions for Array Reading Utility Modules

The array reading utility modules provide a common way for all packages to read variables that have multiple values. The term “array” is simply a programming term for a variable that contains multiple values. There are three modules: U2DREL, U2DINT, and U1DREL. U2DREL reads real two-dimensional variables, U2DINT reads integer two-dimensional variables, and U1DREL reads real one-dimensional variables. All of these modules work similarly. They read one array-control record and, optionally, a data array in a format specified on the array-control record. Several alternate structures for the control record are provided. The original fixed-format control records work as documented in McDonald and Harbaugh (1988), and four free-format versions have been added. The free-format versions are described first because they are easier to use.

FREE-FORMAT CONTROL RECORDS FOR ARRAY READERS:

Values in bold italics are keywords that can be specified as uppercase or lowercase. Each control record is limited to a length of 199 characters.

1. ***CONSTANT*** CNSTNT

All values in the array are set equal to CNSTNT.

2. ***INTERNAL*** CNSTNT FMTIN IPRN

The individual array elements will be read from the same file that contains the control record.

3. ***EXTERNAL*** Nunit CNSTNT FMTIN IPRN

The individual array elements will be read from the file unit number specified by Nunit. The name of the file associated with this file unit must be contained in the name file.

4. ***OPEN/CLOSE*** FNAME CNSTNT FMTIN IPRN

The array will be read from the file whose name is specified by FNAME. This file will be opened on unit 99 just prior to reading the array and closed immediately after the array is read. This file should not be included in the name file. A file that is read using this control record can contain only a single array. The OPEN/CLOSE option is particularly useful for running simulations that require more than 99 files using a computer that allows only 99 files to be opened simultaneously.

FIXED-FORMAT CONTROL RECORD FOR ARRAY READERS:

A fixed-format control record contains the following variables:

LOCAT CNSTNT FMTIN IPRN

These variables are explained below. LOCAT, CNSTNT, and IPRN are 10-character numeric fields. For U2DREL and U1DREL, CNSTNT is a real number. For U2DINT, CNSTNT is an integer and must not include a decimal. FMTIN is a 20-character text field. All four variables are always read when the control record is fixed format; however, some of the variables are unused in some situations. For example when LOCAT = 0, FMTIN and IPRN are not used.

Explanation of Variables in the Array Control Records

CNSTNT—is a real-number constant for U2DREL and U1DREL, and an integer constant for U2DINT. If the array is being defined as a constant, CNSTNT is the constant value. If individual elements of the array are being read, the values are multiplied by CNSTNT after they are read. When CNSTNT is used as a multiplier and specified as 0, it is changed to 1.

FMTIN—is the format for reading array elements. The format must contain 20 characters or less. The format must either be a standard Fortran format that is enclosed in parentheses, "(FREE)" which indicates free format, or "(BINARY)" which indicates binary (unformatted) data. When using a free-format control record, the format must be enclosed in apostrophes if it contains one or more blanks or commas. There are only two ways to create a binary file that can be read by MODFLOW. The first way is to use MODFLOW to create the file by saving heads in a binary file. This is commonly done when it is desired to use computed heads from one simulation as initial heads for a subsequent simulation. The other way to create a binary file is to write a special program that generates a binary file. "(FREE)" and "(BINARY)" can only be specified in free-format control records. Also, "(BINARY)" can only be specified when using U2DREL or U2DINT, and only when the control record is EXTERNAL or OPEN/CLOSE. When the "(FREE)" option is used, be sure that all array elements have a non-blank value and that a comma or at least one blank separates adjacent values.

IPRN—is a flag that indicates if the array being read should be printed (written to the listing file) after it has been read and a code for indicating the format that should be used when it is printed. The format codes are different for each of the three array-reading modules as shown below. IPRN is set to zero when the specified value exceeds those defined. If IPRN is less than zero, the array will not be printed.

IPRN	U2DREL	U2DINT	U1DREL
0	10G11.4	10I11	10G12.5
1	11G10.3	60I1	5G12.5
2	9G13.6	40I2	
3	15F7.1	30I3	
4	15F7.2	25I4	
5	15F7.3	20I5	
6	15F7.4	10I11	
7	20F5.0	25I2	
8	20F5.1	15I4	
9	20F5.2	10I6	
10	20F5.3		
11	20F5.4		
12	10G11.4		
13	10F6.0		
14	10F6.1		
15	10F6.2		
16	10F6.3		
17	10F6.4		
18	10F6.5		
19	5G12.5		
20	6G11.4		
21	7G9.2		

Nunit—is the unit for reading the array when the *EXTERNAL* free-format control record is used.

LOCAT—indicates the location of the array values for a fixed-format array control record. If LOCAT = 0, all elements are set equal to CNSTNT. If LOCAT > 0, it is the unit number for reading formatted records using FMTIN as the format. If LOCAT < 0, it is the unit number for binary (unformatted) records, and FMTIN is ignored. Also, when LOCAT is not 0, the array values are multiplied by CNSTNT after they are read.

Examples of Free-Format Control Records for Reading an Array

The following examples use free-format control records for reading an array. The example array is a real array consisting of 4 rows with 7 columns per row:

CONSTANT 5.7 This sets an entire array to the value "5.7".

INTERNAL 1.0 (7F4.0) 3 This reads the array values from the
 1.2 3.7 9.3 4.2 2.2 9.9 1.0 file that contains the control record.
 3.3 4.9 7.3 7.5 8.2 8.7 6.6 Thus, the values immediately follow the
 4.5 5.7 2.2 1.1 1.7 6.7 6.9 control record.
 7.4 3.5 7.8 8.5 7.4 6.8 8.8

EXTERNAL 52 1.0 (7F4.0) 3 This reads the array from the formatted
 file opened on unit 52.

EXTERNAL 47 1.0 (BINARY) 3 This reads the array from the
 binary file opened on unit 47.

OPEN/CLOSE test.dat 1.0 (7F4.0) 3 This reads the array from the
 file named "test.dat".

Input Instructions for List Utility Module (ULSTRD)

Module ULSTRD reads lists, which are any number of repetitions of an input item that contains multiple variables. Examples of packages that make use of this module are the General-Head Boundary, Drain, River, and Well Packages.

1. [**EXTERNAL** IN] or [**OPEN/CLOSE** FNAME]
 If Item 1 is not included, then the list is read from the package file. Item 1 must begin with the keyword "EXTERNAL" or the keyword "OPEN/CLOSE" (not both).
2. [**SFAC** Scale]
3. List

Explanation of Variables Read by the List Utility Module

IN—is the unit number for a file from which the list will be read. The name of the file associated with this file unit must be contained in the name file, and its file type must be "DATA" in the name file.

FNAME—is the name of a file from which the list will be read. This file will be opened on unit 99 just before reading the list and closed immediately after the list is read. This file should not be included in the name file. The OPEN/CLOSE option is particularly useful for running simulations that require more than 99 files using a computer that allows only 99 files to be opened simultaneously.

Scale—is a scale factor that is multiplied times the value of one or more variables within every record of the list. The input instructions that define a list, which will be read by ULSTRD, should specify the variables to which SFAC applies. If Item 2 is not included, then Scale is 1.0. If Item 2 is included, it must begin with the keyword "SFAC." The values of the list variables that are printed to the listing file include the effect of Scale.

List—is a specified number of lines of data in which each line contains a specified number of variables. The first three variables are always layer, row, and column. The other fields vary according to which package is calling this module.

EXAMPLE

The example problem described in McDonald and Harbaugh (1988) and duplicated in Harbaugh and McDonald (1996a) is used here to demonstrate the use of MODFLOW-2000. The example is implemented two ways—with and without parameters. For the implementation without parameters, the BCF Package is used as in McDonald and Harbaugh (1988). When the example is implemented using parameters, the LPF Package is used. The simulated system is shown in figure 10 (McDonald and Harbaugh, 1988, Appendix D).

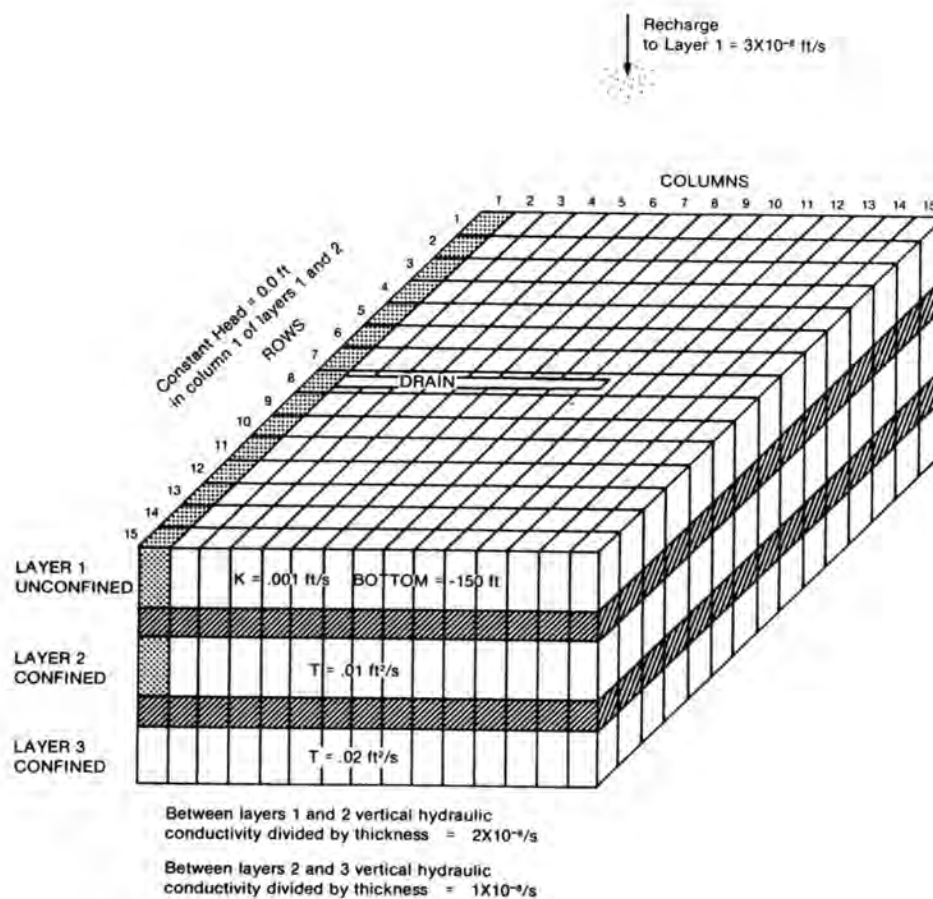


Figure 10. -- Illustration of the system simulated in the example problem.

Example Without Parameters

To run the problem without parameters, the example input data from Harbaugh and McDonald (1996a) was converted by the conversion program described earlier, MF96TO2K. The conversion program requires the user to specify which model layers are underlain by Quasi-3D confining beds and any elevation data that are not included in the BCF Package input file. For this problem, the following elevation information must be provided to the conversion program:

- the elevation of the top of layer 1,
- the elevations of the bottoms of the confining beds under layers 1 and 2, and
- the elevations of the bottoms of model layers 2 and 3.

As shown in figure 10, all the layers are flat, but there are no elevations specified except for the bottom of layer 1. The other elevations were not specified in the original problem because they were not needed in earlier versions of MODFLOW. Layer thickness is incorporated into transmissivity and vertical leakance (VCONT) values. For MODFLOW-2000, however, all of the elevations must be specified. Any physically consistent elevations can be used if the only goal is to reproduce the heads calculated by MODFLOW-96. That is, each successive layer must have a lower elevation than the layer above, and the top elevation of layer 1 must be above the water table. Within these constraints, the following values were arbitrarily selected:

Elevation of the top of layer 1 = 200 ft,
 Elevation of the bottom of the confining bed below layer 1 = -200 ft,
 Elevation of the bottom of layer 2 = -300 ft,
 Elevation of the bottom of the confining bed below layer 2 = -350 ft, and
 Elevation of the bottom of layer 3 = -450 ft.

The following are the resulting input files:

Name File:

```
LIST 6 twri.lst
BAS6      5 TWRI.ba6
BCF6      11 TWRI.bc6
WEL 12 twri.wel
DRN 13 twri.drn
RCH 18 twri.rch
SIP 19 twri.sip
OC 22 twri.oc
DIS      10 TWRI.dis
```

Discretization (DIS) File

```
3 15 15 1 1 0 NLAY,NROW,NCOL,NPER,ITMUNI,LENUNI
1 1 0
CONSTANT 5.000000E+03 DELR
CONSTANT 5.000000E+03 DELC
CONSTANT 2.000000E+02 TOP of system
CONSTANT -1.500000E+02 Layer BOTM layer 1
CONSTANT -2.000000E+02 Confining bed BOTM layer 1
CONSTANT -3.000000E+02 Layer BOTM layer 2
CONSTANT -3.500000E+02 Confining bed BOTM layer 2
CONSTANT -4.500000E+02 Layer BOTM layer 3
8.640E+04 1 1.000E+00 SS PERLEN,NSTP,TSMULT,Ss/tr
```

Basic Package (BAS6) File

```
#SAMPLE---3 LAYERS, 15 ROWS, 15 COLUMNS; STEADY STATE; CONSTANT HEADS COLUMN 1
#LAYERS 1 AND 2; RECHARGE, WELLS AND DRAINS
NO OPTIONS
```

```
INTERNAL          1 (2014)      3 IBOUND layer  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
INTERNAL          1 (2014)      3 IBOUND layer  2
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
-1  1  1  1  1  1  1  1  1  1  1  1  1  1  1
CONSTANT          1 IBOUND layer  3
  999.99  HNOFLO
CONSTANT  0.000000E+00  Initial Head layer  1
CONSTANT  0.000000E+00  Initial Head layer  2
CONSTANT  0.000000E+00  Initial Head layer  3
```

Output Control Option (OC) File—Default output control was used when this problem was run by McDonald and Harbaugh (1988), so no file was required. The Output Control Option is used here only for the purpose of changing the format for printing head. A format has been chosen that uses less page width.

```
HEAD PRINT FORMAT 20
PERIOD 1 STEP 1
PRINT HEAD
```

Block-Centered Flow Package (BCF6) File

```

0 1.00E+30      0 0.00E+00      0      0
1 0 0
CONSTANT 1.000000E+00 TRPY
CONSTANT 1.000000E-03 HY layer 1
CONSTANT 2.000000E-08 VCONT layer 1
CONSTANT 1.000000E-02 TRAN layer 2
CONSTANT 1.000000E-08 VCONT layer 2
CONSTANT 2.000000E-02 TRAN layer 3

```

Recharge Package (RCH) File

```

1      0 NRCHOP, IRCHBD
1      INRECH
0      3.E-8 RECH-1

```

Well Package (WEL) File

```

15      0 MXACTW, IWELCB
15      ITMP
3      5      11      -5.
2      4      6      -5.
2      6      12      -5.
1      9      8      -5.
1      9      10     -5.
1      9      12     -5.
1      9      14     -5.
1      11     8      -5.
1      11     10     -5.
1      11     12     -5.
1      11     14     -5.
1      13     8      -5.
1      13     10     -5.
1      13     12     -5.
1      13     14     -5.

```

Drain Package (DRN) File

```

9      0 MXACTD, IDRNCB
9      ITMP
1      8      2      0.      1.E00
1      8      3      0.      1.E00
1      8      4      10.     1.E00
1      8      5      20.     1.E00
1      8      6      30.     1.E00
1      8      7      50.     1.E00
1      8      8      70.     1.E00
1      8      9      90.     1.E00
1      8      10     100.    1.E00

```

Strongly Implicit Procedure Package (SIP) File

```
50          5          MXITER, NPARM
1.         .001          0          .001          1
```

This problem was run on a personal computer using a runfile that was compiled using the Lahey Fortran 95 compiler. The Name File includes only a LIST output file (i.e., there is no GLOBAL file), so all of the “printed” output is in a single file. The contents of the LIST file are shown in figure 11.

```
MODFLOW-2000
U.S. GEOLOGICAL SURVEY MODULAR FINITE-DIFFERENCE GROUND-WATER FLOW MODEL
VERSION 1.00 06/13/2000
```

This model run combines GLOBAL and LIST output into this single file.

```
GLOBAL LISTING FILE: twri.lst
                        UNIT    6

OPENING TWRI.ba6
FILE TYPE:BAS6  UNIT    5

OPENING TWRI.bc6
FILE TYPE:BCF6  UNIT   11

OPENING twri.wel
FILE TYPE:WEL   UNIT   12

OPENING twri.drn
FILE TYPE:DRN   UNIT   13

OPENING twri.rch
FILE TYPE:RCH   UNIT   18

OPENING twri.sip
FILE TYPE:SIP   UNIT   19

OPENING twri.oc
FILE TYPE:OC    UNIT   22

OPENING TWRI.dis
FILE TYPE:DIS   UNIT   10

DISCRETIZATION INPUT DATA READ FROM UNIT 10
  3 LAYERS      15 ROWS      15 COLUMNS
  1 STRESS PERIOD(S) IN SIMULATION
MODEL TIME UNIT IS SECONDS
MODEL LENGTH UNIT IS UNDEFINED
```

Figure 11. -- LIST file for example problem without parameters.

THE OBSERVATION PROCESS IS INACTIVE
 THE SENSITIVITY PROCESS IS INACTIVE
 THE PARAMETER-ESTIMATION PROCESS IS INACTIVE

MODE: FORWARD

Confining bed flag for each layer:

1 1 0

6555 ELEMENTS OF GX ARRAY USED OUT OF 6555
 675 ELEMENTS OF GZ ARRAY USED OUT OF 675
 675 ELEMENTS OF IG ARRAY USED OUT OF 675

DELR = 5000.00

DELC = 5000.00

TOP ELEVATION OF LAYER 1 = 200.000

MODEL LAYER BOTTOM EL. = -150.000 FOR LAYER 1

BOT. EL. OF QUASI-3D BED = -200.000 FOR LAYER 1

MODEL LAYER BOTTOM EL. = -300.000 FOR LAYER 2

BOT. EL. OF QUASI-3D BED = -350.000 FOR LAYER 2

MODEL LAYER BOTTOM EL. = -450.000 FOR LAYER 3

STRESS PERIOD	LENGTH	TIME STEPS	MULTIPLIER FOR DELT	SS FLAG
1	86400.00	1	1.000	SS

STEADY-STATE SIMULATION

SIP5 -- STRONGLY IMPLICIT PROCEDURE SOLUTION PACKAGE
 VERSION 5, 9/1/93 INPUT READ FROM UNIT 19
 MAXIMUM OF 50 ITERATIONS ALLOWED FOR CLOSURE
 5 ITERATION PARAMETERS
 2755 ELEMENTS IN X ARRAY ARE USED BY SIP
 150 ELEMENTS IN IX ARRAY ARE USED BY SIP

2755 ELEMENTS OF X ARRAY USED OUT OF 2755
 0 ELEMENTS OF Z ARRAY USED OUT OF 1
 150 ELEMENTS OF IX ARRAY USED OUT OF 150
 0 ELEMENTS OF XHS ARRAY USED OUT OF 1

Figure 11. -- LIST file for example problem without parameters—Continued.

SOLUTION BY THE STRONGLY IMPLICIT PROCEDURE

```

-----
MAXIMUM ITERATIONS ALLOWED FOR CLOSURE =      50
      ACCELERATION PARAMETER =      1.0000
HEAD CHANGE CRITERION FOR CLOSURE =    0.10000E-02
SIP HEAD CHANGE PRINTOUT INTERVAL =      1

```

5 ITERATION PARAMETERS CALCULATED FROM SPECIFIED WSEED = 0.00100000 :

```
0.000000E+00 0.822172E+00 0.968377E+00 0.994377E+00 0.999000E+00
```

```

#SAMPLE----3 LAYERS, 15 ROWS, 15 COLUMNS; STEADY STATE; CONSTANT HEADS COLUMN 1
#LAYERS 1 AND 2; RECHARGE, WELLS AND DRAINS
  3 LAYERS      15 ROWS      15 COLUMNS
  1 STRESS PERIOD(S) IN SIMULATION

```

```

BAS6 -- BASIC PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 5
      15 ELEMENTS IN IR ARRAY ARE USED BY BAS

```

```

BCF6 -- BLOCK-CENTERED FLOW PACKAGE, VERSION 6, 1/11/2000
      INPUT READ FROM UNIT 11

```

STEADY-STATE SIMULATION

HEAD AT CELLS THAT CONVERT TO DRY= 0.10000E+31

WETTING CAPABILITY IS NOT ACTIVE

LAYER	LAYER-TYPE CODE	INTERBLOCK T
1	1	0 -- HARMONIC
2	0	0 -- HARMONIC
3	0	0 -- HARMONIC

678 ELEMENTS IN RX ARRAY ARE USED BY BCF

```

WEL6 -- WELL PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 12

```

No named parameters

MAXIMUM OF 15 ACTIVE WELLS AT ONE TIME

60 ELEMENTS IN RX ARRAY ARE USED BY WEL

```

DRN6 -- DRAIN PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 13

```

No named parameters

MAXIMUM OF 9 ACTIVE DRAINS AT ONE TIME

45 ELEMENTS IN RX ARRAY ARE USED BY DRN

```

RCH6 -- RECHARGE PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 18

```

No named parameters

OPTION 1 -- RECHARGE TO TOP LAYER

225 ELEMENTS IN RX ARRAY ARE USED BY RCH

225 ELEMENTS IN IR ARRAY ARE USED BY RCH

1008 ELEMENTS OF RX ARRAY USED OUT OF 1008

240 ELEMENTS OF IR ARRAY USED OUT OF 240

1

```

#SAMPLE----3 LAYERS, 15 ROWS, 15 COLUMNS; STEADY STATE; CONSTANT HEADS COLUMN 1
#LAYERS 1 AND 2; RECHARGE, WELLS AND DRAINS

```

Figure 11. -- LIST file for example problem without parameters—Continued.


```

          BOUNDARY ARRAY FOR LAYER 1
READING ON UNIT 5 WITH FORMAT: (20I4)

      1  2  3  4  5  6  7  8  9 10 11 12 13 14 15
.....
1  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
2  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
3  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
4  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
5  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
6  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
7  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
8  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
9  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
10 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
11 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
12 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
13 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
14 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
15 -1  1  1  1  1  1  1  1  1  1  1  1  1  1

```

```

          BOUNDARY ARRAY FOR LAYER 2
READING ON UNIT 5 WITH FORMAT: (20I4)

      1  2  3  4  5  6  7  8  9 10 11 12 13 14 15
.....
1  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
2  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
3  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
4  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
5  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
6  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
7  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
8  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
9  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
10 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
11 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
12 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
13 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
14 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
15 -1  1  1  1  1  1  1  1  1  1  1  1  1  1

```

BOUNDARY ARRAY = 1 FOR LAYER 3

AQUIFER HEAD WILL BE SET TO 999.99 AT ALL NO-FLOW NODES (IBOUND=0).

INITIAL HEAD = 0.000000 FOR LAYER 1

INITIAL HEAD = 0.000000 FOR LAYER 2

INITIAL HEAD = 0.000000 FOR LAYER 3

Figure 11. -- LIST file for example problem without parameters—Continued.

OUTPUT CONTROL IS SPECIFIED ONLY AT TIME STEPS FOR WHICH OUTPUT IS DESIRED
 HEAD PRINT FORMAT CODE IS 20 DRAWDOWN PRINT FORMAT CODE IS 0
 HEADS WILL BE SAVED ON UNIT 0 DRAWDOWNS WILL BE SAVED ON UNIT 0

COLUMN TO ROW ANISOTROPY = 1.00000

HYD. COND. ALONG ROWS = 1.000000E-03 FOR LAYER 1

VERT HYD COND /THICKNESS = 2.000000E-08 FOR LAYER 1

TRANSMIS. ALONG ROWS = 1.000000E-02 FOR LAYER 2

VERT HYD COND /THICKNESS = 1.000000E-08 FOR LAYER 2

TRANSMIS. ALONG ROWS = 2.000000E-02 FOR LAYER 3

0 Well parameters

0 Drain parameters

0 Recharge parameters

1
 STRESS PERIOD NO. 1, LENGTH = 86400.00

NUMBER OF TIME STEPS = 1

MULTIPLIER FOR DELT = 1.000

INITIAL TIME STEP SIZE = 86400.00

WELL NO.	LAYER	ROW	COL	STRESS RATE
1	3	5	11	-5.000
2	2	4	6	-5.000
3	2	6	12	-5.000
4	1	9	8	-5.000
5	1	9	10	-5.000
6	1	9	12	-5.000
7	1	9	14	-5.000
8	1	11	8	-5.000
9	1	11	10	-5.000
10	1	11	12	-5.000
11	1	11	14	-5.000
12	1	13	8	-5.000
13	1	13	10	-5.000
14	1	13	12	-5.000
15	1	13	14	-5.000

15 WELLS

Figure 11. -- LIST file for example problem without parameters—Continued.

DRAIN NO.	LAYER	ROW	COL	DRAIN EL.	CONDUCTANCE
1	1	8	2	0.0000	1.000
2	1	8	3	0.0000	1.000
3	1	8	4	10.00	1.000
4	1	8	5	20.00	1.000
5	1	8	6	30.00	1.000
6	1	8	7	50.00	1.000
7	1	8	8	70.00	1.000
8	1	8	9	90.00	1.000
9	1	8	10	100.0	1.000

9 DRAINS

RECHARGE = 3.000000E-08

SOLVING FOR HEAD

31 ITERATIONS FOR TIME STEP 1 IN STRESS PERIOD 1

MAXIMUM HEAD CHANGE FOR EACH ITERATION:

HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL
-22.41 (3, 5, 11)	12.48 (1, 1, 15)	13.39 (3, 1, 14)	48.21 (1, 1, 15)	35.91 (3, 1, 13)
2.482 (1, 9, 14)	1.430 (3, 10, 13)	6.214 (1, 12, 14)	7.411 (3, 11, 14)	13.66 (1, 15, 15)
0.5503 (3, 8, 7)	0.4821 (2, 6, 9)	0.4711 (3, 5, 10)	2.019 (1, 11, 14)	2.302 (3, 5, 13)
0.1108 (1, 13, 12)	0.7059E-01 (3, 12, 11)	0.2819 (1, 14, 14)	0.3141 (3, 13, 14)	0.3320 (1, 15, 15)
0.7853E-02 (1, 13, 12)	0.1586E-01 (2, 11, 11)	0.1777E-01 (3, 11, 10)	0.7910E-01 (1, 14, 14)	0.8500E-01 (3, 7, 14)
0.4169E-02 (1, 13, 14)	0.2555E-02 (3, 14, 15)	0.9769E-02 (1, 14, 14)	0.1082E-01 (3, 13, 14)	0.1030E-01 (1, 15, 15)
0.2430E-03 (1, 13, 12)				

OUTPUT CONTROL FOR STRESS PERIOD 1 TIME STEP 1
PRINT HEAD FOR ALL LAYERS

1

HEAD IN LAYER 1 AT END OF TIME STEP 1 IN STRESS PERIOD 1

	1	2	3	4	5	6
	7	8	9	10	11	12
	13	14	15			
1	0.0000	24.94	44.01	59.26	71.82	82.52
	91.91	100.0	106.9	112.6	117.4	121.3
	124.3	126.4	127.4			

Figure 11. -- LIST file for example problem without parameters—Continued.

2	0.0000	24.45	43.10	57.98	70.17	80.57
	90.12	98.40	105.3	111.0	115.7	119.6
	122.7	124.9	126.1			
3	0.0000	23.45	41.30	55.43	66.78	76.21
	86.51	95.20	102.2	107.6	112.0	116.1
	119.6	122.1	123.4			
4	0.0000	21.92	38.61	51.75	61.79	68.03
	81.34	90.75	97.64	102.5	106.1	110.7
	114.9	117.9	119.4			
5	0.0000	19.73	34.92	47.32	57.69	66.74
	77.09	85.76	92.22	96.15	97.29	103.1
	108.8	112.5	114.3			
6	0.0000	16.51	29.50	40.90	51.30	61.21
	71.19	79.85	86.47	90.82	93.03	94.23
	102.1	106.4	108.4			
7	0.0000	11.55	21.10	31.21	41.40	51.84
	63.08	72.68	79.95	84.92	88.60	91.66
	96.43	99.82	101.8			
8	0.0000	3.483	6.832	16.25	26.30	36.97
	52.59	64.31	72.52	77.25	81.99	85.00
	89.27	91.72	94.33			
9	0.0000	10.54	19.11	28.12	36.92	45.27
	52.95	55.38	65.15	66.07	73.93	73.79
	80.84	80.17	86.49			
10	0.0000	14.62	25.86	35.38	43.49	50.11
	54.93	57.55	62.95	65.55	70.39	72.44
	76.72	78.26	81.79			
11	0.0000	17.11	29.96	40.01	47.78	53.24
	55.81	53.33	60.27	59.29	66.43	65.45
	72.22	71.04	77.62			
12	0.0000	18.68	32.56	43.07	50.81	55.92
	58.33	58.47	61.93	63.18	67.12	68.50
	72.29	73.46	76.85			
13	0.0000	19.67	34.24	45.14	53.01	58.04
	59.91	56.75	62.59	60.91	67.22	65.75
	71.90	70.35	76.48			
14	0.0000	20.27	35.27	46.48	54.61	60.08
	63.17	64.52	67.25	68.79	71.64	73.18
	75.84	77.03	79.09			
15	0.0000	20.56	35.78	47.16	55.48	61.26
	65.02	67.52	69.94	72.01	74.29	76.22
	78.22	79.66	80.82			

1

HEAD IN LAYER 2 AT END OF TIME STEP 1 IN STRESS PERIOD 1

	1	2	3	4	5	6
	7	8	9	10	11	12
	13	14	15			
1	0.0000	24.66	43.73	59.02	71.61	82.32
	91.72	99.86	106.7	112.5	117.2	121.1
	124.1	126.2	127.3			

.....

Figure 11. -- LIST file for example problem without parameters—Continued.

2	0.0000	24.17	42.83	57.74	69.95	80.36
	89.93	98.22	105.1	110.8	115.5	119.4
	122.6	124.8	125.9			
3	0.0000	23.17	41.03	55.19	66.53	75.77
	86.29	95.02	102.0	107.4	111.8	116.0
	119.5	121.9	123.2			
4	0.0000	21.65	38.34	51.50	61.35	60.17
	80.90	90.55	97.45	102.3	105.4	110.4
	114.8	117.7	119.2			
5	0.0000	19.48	34.65	47.07	57.44	66.30
	76.85	85.57	92.00	95.41	91.09	102.1
	108.6	112.4	114.2			
6	0.0000	16.27	29.24	40.65	51.07	60.98
	70.98	79.65	86.28	90.54	92.06	86.23
	101.7	106.2	108.3			
7	0.0000	11.38	20.95	31.05	41.25	51.70
	62.90	72.48	79.76	84.73	88.35	91.24
	96.22	99.65	101.6			
8	0.0000	4.209	8.330	17.58	27.58	38.25
	52.94	64.19	72.34	77.12	81.81	84.86
	89.10	91.59	94.17			
9	0.0000	10.38	18.96	27.98	36.79	45.16
	52.86	56.13	65.08	66.79	73.87	74.48
	80.77	80.84	86.38			
10	0.0000	14.40	25.61	35.15	43.27	49.91
	54.76	57.48	62.79	65.49	70.24	72.37
	76.57	78.20	81.64			
11	0.0000	16.87	29.70	39.78	47.56	53.05
	55.68	54.09	60.20	60.04	66.37	66.18
	72.16	71.75	77.51			
12	0.0000	18.43	32.31	42.85	50.60	55.73
	58.16	58.41	61.78	63.12	66.98	68.44
	72.15	73.40	76.69			
13	0.0000	19.42	33.98	44.91	52.80	57.85
	59.78	57.50	62.53	61.65	67.16	66.48
	71.84	71.06	76.37			
14	0.0000	20.02	35.02	46.26	54.41	59.88
	62.99	64.39	67.08	68.66	71.48	73.06
	75.68	76.91	78.93			
15	0.0000	20.30	35.52	46.94	55.28	61.07
	64.84	67.34	69.76	71.84	74.11	76.04
	78.04	79.49	80.65			

1

HEAD IN LAYER 3 AT END OF TIME STEP 1 IN STRESS PERIOD 1

	1	2	3	4	5	6
	7	8	9	10	11	12
	13	14	15			
1	1.800	24.34	43.36	58.70	71.33	82.06
	91.48	99.63	106.5	112.3	117.0	120.9
	123.9	126.0	127.1			

.....

Figure 11. -- LIST file for example problem without parameters—Continued.

2	1.764	23.85	42.46	57.42	69.66	80.07
	89.68	97.99	104.9	110.6	115.3	119.2
	122.4	124.6	125.7			
3	1.691	22.86	40.67	54.87	66.20	75.28
	85.98	94.77	101.7	107.2	111.5	115.7
	119.3	121.7	123.0			
4	1.578	21.35	37.98	51.17	60.85	62.69
	80.41	90.28	97.19	101.9	104.1	110.0
	114.5	117.5	119.0			
5	1.415	19.18	34.30	46.75	57.10	65.80
	76.54	85.30	91.67	94.17	77.46	100.7
	108.2	112.1	114.0			
6	1.176	15.99	28.91	40.33	50.76	60.67
	70.70	79.38	86.01	90.12	90.60	88.55
	101.2	106.0	108.0			
7	0.8273	11.21	20.79	30.88	41.09	51.55
	62.67	72.22	79.50	84.46	87.98	90.77
	95.94	99.41	101.4			
8	0.4331	5.131	10.19	19.27	29.19	39.84
	53.40	64.07	72.11	76.95	81.58	84.68
	88.88	91.44	93.95			
9	0.7543	10.22	18.82	27.84	36.66	45.06
	52.78	57.03	65.02	67.64	73.81	75.31
	80.72	81.64	86.24			
10	1.039	14.13	25.29	34.85	42.99	49.65
	54.54	57.44	62.61	65.44	70.05	72.33
	76.39	78.15	81.43			
11	1.224	16.59	29.37	39.47	47.28	52.79
	55.53	55.01	60.16	60.94	66.33	67.06
	72.13	72.60	77.38			
12	1.341	18.15	31.97	42.54	50.32	55.47
	57.94	58.37	61.60	63.08	66.80	68.41
	71.97	73.36	76.49			
13	1.415	19.14	33.65	44.61	52.53	57.60
	59.63	58.39	62.48	62.54	67.12	67.35
	71.80	71.90	76.24			
14	1.460	19.73	34.68	45.96	54.13	59.63
	62.76	64.24	66.87	68.52	71.27	72.91
	75.47	76.77	78.71			
15	1.481	20.01	35.18	46.63	55.00	60.81
	64.59	67.11	69.52	71.61	73.87	75.82
	77.81	79.27	80.42			

1
VOLUMETRIC BUDGET FOR ENTIRE MODEL AT END OF TIME STEP 1 IN STRESS PERIOD 1

CUMULATIVE VOLUMES	L**3	RATES FOR THIS TIME STEP	L**3/T
-----		-----	

Figure 11. -- LIST file for example problem without parameters—Continued.

IN:		IN:	
---		---	
STORAGE =	0.0000	STORAGE =	0.0000
CONSTANT HEAD =	0.0000	CONSTANT HEAD =	0.0000
WELLS =	0.0000	WELLS =	0.0000
DRAINS =	0.0000	DRAINS =	0.0000
RECHARGE =	13608000.0000	RECHARGE =	157.5000
TOTAL IN =	13608000.0000	TOTAL IN =	157.5000
OUT:		OUT:	
----		----	
STORAGE =	0.0000	STORAGE =	0.0000
CONSTANT HEAD =	4326522.5000	CONSTANT HEAD =	50.0755
WELLS =	6480000.0000	WELLS =	75.0000
DRAINS =	2801081.2500	DRAINS =	32.4199
RECHARGE =	0.0000	RECHARGE =	0.0000
TOTAL OUT =	13607603.0000	TOTAL OUT =	157.4954
IN - OUT =	397.0000	IN - OUT =	4.5929E-03
PERCENT DISCREPANCY =	0.00	PERCENT DISCREPANCY =	0.00

	TIME SUMMARY AT END OF TIME STEP 1 IN STRESS PERIOD 1				
	SECONDS	MINUTES	HOURS	DAYS	YEARS

TIME STEP LENGTH	86400.	1440.0	24.000	1.0000	2.73785E-03
STRESS PERIOD TIME	86400.	1440.0	24.000	1.0000	2.73785E-03
TOTAL TIME	86400.	1440.0	24.000	1.0000	2.73785E-03

1

Figure 11. -- LIST file for example problem without parameters—Continued.

Example With Parameters

The same problem is repeated using the LPF Package. LPF does not have an option to enter transmissivity for confined layers; instead, horizontal hydraulic conductivity (HK) must be entered for all layers. HK values for layers 2 and 3 were not directly specified in the problem, but HK can be easily calculated from the specified values of transmissivity and the assumed layer thicknesses.

LPF also differs from BCF in the way vertical hydraulic information is entered. LPF requires vertical hydraulic conductivity information for each model layer and Quasi-3D confining bed, whereas BCF requires only the vertical leakage (VCONT) between nodes, which is the average vertical hydraulic conductivity (including a Quasi-3D confining bed if it exists) divided by the distance between nodes. The BCF user must calculate a representative VCONT from the VCONT of three individual components. For example, for a cell in layer 1, VCONT is calculated from the VCONT of half the thickness of layer 1, the full thickness of the confining bed, and half the thickness of layer 2 (McDonald and Harbaugh, 1988, p. 5-16). There is no unique way to calculate the vertical hydraulic conductivity of each layer based on the BCF VCONT values, which is all that the original

problem specified. The approach used here is to assume that the confining bed dominates the VCONT calculation and that there is virtually no contribution from the actual model layers, which is consistent with the assumptions used in making the Quasi-3D approximation. If the model layers have no influence on VCONT, then VKCB is VCONT times the confining-bed thickness. For layer 1, VKCB is 10^{-6} ft/s; for layer 2 it is 5×10^{-7} ft/s.

Although it is assumed for the above calculation of VKCB that there is no contribution to CV from the aquifer vertical hydraulic conductivity (VK), LPF requires VK to be specified as part of the input data. According to equation 27, VK should be infinite in order to have no impact on CV. There is no way to directly specify an infinite value for VK, but the effect of having an infinite value can be obtained by making the VK values large compared to VKCB. For this example problem, VK is set to 1.0 ft/s for all model layers, which is a million times larger than the largest VKCB. The low impact of this value on CV is demonstrated by the fact that the calculated head as shown in the LIST file when LPF is used (fig. 13) is the same as the head in the BCF LIST file (fig. 11).

Note that any combination of VK and CBVK that produces virtually the same CV that BCF calculates using the original VCONT values will produce the same heads in this problem. If the values of VK for layer 1 were relatively small compared to CBVK, then the VK values would have to be varied from cell to cell due to the fact that the saturated thickness of layer 1 varies; that is, as indicated by equation 27, the conductance between cells depends on saturated thickness of the model cells. Thus, the choice to use a large VK in order to essentially eliminate the aquifer contribution to CV makes the LPF input simpler than it would otherwise be.

Another difference in the input data for this problem is that parameters are used for defining some of the data in order to illustrate the use of parameters. Parameters are used for defining HK, VKCB, RECH, WELL, and DRAI input variables. In the LPF Package, five parameters are used—three define HK, and two define VKCB. Each of these parameters is for a single layer, and each parameter has a uniform value for the entire layer. HK1, HK2, and HK3 are HK for layers 1, 2, and 3, respectively. These use “NONE” for the multiplier array, which means that there is no multiplier array. Instead, the parameter value applies directly to all of the included cells. The zone array is “ALL”, which means the values for all cells in the indicated layer are defined by the parameter. Parameter VKCB1 defines VKCB values for all of layer 1, and VKCB2 defines VKCB values for layer 2. Both of these parameters use the same multiplier array, which has a value of $1.0E-6$ at all cells. The parameter value for VKCB1 is 1.0, and the value for VKCB2 is 0.5, which produces the required values of $1.0E-6$ for layer 1 and $5.0E-7$ for layer 2.

For the RCH Package, two parameters, RCH1 and RCH2, are used. RCH1 includes the left seven columns of the grid, and RCH2 includes the right 8 columns. These areas are specified using zone array RCHZONES. RCHZONES contains “1” in columns 1-7, and “2” in columns 8-15. Accordingly, the zone value of 1 is specified in the definition of RCH1, and zone value 2 is specified for RCH2. The parameter values for RCH1 and RCH2 are both $3.0E-8$, which results in all cells having the same recharge rate. Thus, there is no advantage of having two parameters in this example; however, two zones were used to illustrate how this is done. The value of having two zones would become apparent if there were a need to change the recharge rate in the area of zone 1 without changing the zone 2 values. This could be done simply by changing the value of parameter RCH1.

The Well Package has one parameter, WELL1, and the Drain Package has one parameter, DRN1. Both of these parameters have a value of 1.0, so the multipliers in the lists for both parameters are simply the same as the data values that are used to define the input data without using parameters. There could be an advantage of using parameters, however, if there were a need to change all of the input values associated with the parameters. All of the values could be changed by a constant factor simply by changing the parameter values. The parameters define only part of the wells and drains. Some of them are defined as separate non-parameter lists.

The input files that are different as a result of using LPF instead of BCF and adding parameters are shown below. The DIS, BAS, SIP, and OC files are not shown again because they are unchanged.

Name File		
GLOBAL	1	twri.glo
LIST	6	twri.lst
BAS6	5	TWRI.ba6


```

LPF    11 twri.lpf
WEL    12 twri.wel
DRN    13 twri.drn
RCH    18 twri.rch
SIP    19 twri.sip
OC     22 twri.oc
MULT   8 twri.mlt
ZONE   9 twri.zon
DIS    10 TWRI.dis

```

Multiplier File

```

1
MULT1
CONSTANT 1.0E-6

```

Zone File

```

1
RCHZONES
INTERNAL 1 (15I2) 1
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2

```

Layer-Property Flow Package (LPF) File

```

# LPF Package input data for example problem
0 1.00E+30 5 ILPFCB,HDRY,NPLPF
1 0 0
0 0 0
1. 1. 1.
0 0 0

```

```

0 0 0
HK1 HK 1.0E-3 1
1 NONE ALL
HK2 HK 1.0E-4 1
2 NONE ALL
HK3 HK 2.0E-4 1
3 NONE ALL
VKCB1 VKCB 1.0 1
1 MULT1 ALL
VKCB2 VKCB 0.5 1
2 MULT1 ALL
0 HK layer 1
CONSTANT 1.0 VK layer 1
0 VKCB layer 1
0 HK layer 2
CONSTANT 1.0 VK layer 2
0 VKCB layer 2
0 HK layer 3
CONSTANT 1.0 VK layer 3

```

Recharge Package (RCH) File

```

PARAMETER 2
1 0 NRCHOP,IRCHBD
RCH1 RCH 3.0E-8 1
NONE RCHZONES 1
RCH2 RCH 3.0E-8 1
NONE RCHZONES 2
2 INRECH
RCH1
RCH2

```

Well Package (WEL) File

```

PARAMETER 1 12
15 0 MXACTW,IWELCB
WELL1 Q 1.0 12
1 9 8 -5.
1 9 10 -5.
1 9 12 -5.
1 9 14 -5.
1 11 8 -5.
1 11 10 -5.
1 11 12 -5.
1 11 14 -5.

```

1	13	8	-5.
1	13	10	-5.
1	13	12	-5.
1	13	14	-5.
3	1		ITMP , NP
3	5	11	-5.
2	4	6	-5.
2	6	12	-5.

WELL1

Drain Package (DRN) File

PARAMETER	1	2				
	9	0		MXACTD , IDRNCB		
DRN1	DRN	1.0	2			
	1	8	2	0.	1.E00	
	1	8	3	0.	1.E00	
	7	1		ITMP , NP		
	1	8	4	10.	1.E00	
	1	8	5	20.	1.E00	
	1	8	6	30.	1.E00	
	1	8	7	50.	1.E00	
	1	8	8	70.	1.E00	
	1	8	9	90.	1.E00	
	1	8	10	100.	1.E00	

DRN1

The Name File specifies both GLOBAL and LIST output files. The GLOBAL file is shown in figure 12, and the LIST file is shown in figure 13.

MODFLOW-2000
U.S. GEOLOGICAL SURVEY MODULAR FINITE-DIFFERENCE GROUND-WATER FLOW MODEL
VERSION 1.00 06/13/2000

This model run produced both GLOBAL and LIST files. This is the GLOBAL file.

```
GLOBAL LISTING FILE: twri.glo
                        UNIT  1

OPENING twri.lst
FILE TYPE:LIST  UNIT  6

OPENING TWRI.ba6
FILE TYPE:BAS6  UNIT  5

OPENING twri.lpf
FILE TYPE:LPF   UNIT 11

OPENING twri.wel
FILE TYPE:WEL   UNIT 12

OPENING twri.drn
FILE TYPE:DRN   UNIT 13

OPENING twri.rch
FILE TYPE:RCH   UNIT 18

OPENING twri.sip
FILE TYPE:SIP   UNIT 19

OPENING twri.oc
FILE TYPE:OC    UNIT 22

OPENING twri.mlt
FILE TYPE:MULT  UNIT  8

OPENING twri.zon
FILE TYPE:ZONE  UNIT  9

OPENING TWRI.dis
FILE TYPE:DIS   UNIT 10

DISCRETIZATION INPUT DATA READ FROM UNIT 10
  3 LAYERS      15 ROWS      15 COLUMNS
  1 STRESS PERIOD(S) IN SIMULATION
MODEL TIME UNIT IS SECONDS
MODEL LENGTH UNIT IS UNDEFINED
```

Figure 12. -- GLOBAL file for example problem with parameters.

THE OBSERVATION PROCESS IS INACTIVE
THE SENSITIVITY PROCESS IS INACTIVE
THE PARAMETER-ESTIMATION PROCESS IS INACTIVE

MODE: FORWARD

ZONE OPTION, INPUT READ FROM UNIT 9
1 ZONE ARRAYS

MULTIPLIER OPTION, INPUT READ FROM UNIT 8
1 MULTIPLIER ARRAYS

Confining bed flag for each layer:

1 1 0

6780 ELEMENTS OF GX ARRAY USED OUT OF 6780
675 ELEMENTS OF GZ ARRAY USED OUT OF 675
900 ELEMENTS OF IG ARRAY USED OUT OF 900

DELR = 5000.00

DELC = 5000.00

TOP ELEVATION OF LAYER 1 = 200.000

MODEL LAYER BOTTOM EL. = -150.000 FOR LAYER 1

BOT. EL. OF QUASI-3D BED = -200.000 FOR LAYER 1

MODEL LAYER BOTTOM EL. = -300.000 FOR LAYER 2

BOT. EL. OF QUASI-3D BED = -350.000 FOR LAYER 2

MODEL LAYER BOTTOM EL. = -450.000 FOR LAYER 3

STRESS PERIOD	LENGTH	TIME STEPS	MULTIPLIER FOR DELT	SS FLAG
1	86400.00	1	1.000	SS

STEADY-STATE SIMULATION

MULT. ARRAY: MULT1 = 1.000000E-06

ZONE ARRAY: RCHZONES
READING ON UNIT 9 WITH FORMAT: (15I2)

Figure 12. -- GLOBAL file for example problem with parameters—Continued.

```

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15
.....
1 1 1 1 1 1 1 2 2 2 2 2 2 2 2
2 1 1 1 1 1 1 2 2 2 2 2 2 2 2
3 1 1 1 1 1 1 2 2 2 2 2 2 2 2
4 1 1 1 1 1 1 2 2 2 2 2 2 2 2
5 1 1 1 1 1 1 2 2 2 2 2 2 2 2
6 1 1 1 1 1 1 2 2 2 2 2 2 2 2
7 1 1 1 1 1 1 2 2 2 2 2 2 2 2
8 1 1 1 1 1 1 2 2 2 2 2 2 2 2
9 1 1 1 1 1 1 2 2 2 2 2 2 2 2
10 1 1 1 1 1 1 2 2 2 2 2 2 2 2
11 1 1 1 1 1 1 2 2 2 2 2 2 2 2
12 1 1 1 1 1 1 2 2 2 2 2 2 2 2
13 1 1 1 1 1 1 2 2 2 2 2 2 2 2
14 1 1 1 1 1 1 2 2 2 2 2 2 2 2
15 1 1 1 1 1 1 2 2 2 2 2 2 2 2

```

```

LPF1 -- LAYER PROPERTY FLOW PACKAGE, VERSION 1, 1/11/2000
      INPUT READ FROM UNIT 11
# LPF Package input data for example problem
HEAD AT CELLS THAT CONVERT TO DRY= 1.00000E+30
      5 Named Parameters

```

LAYER FLAGS:

LAYER	LAYTYP	LAYAVG	CHANI	LAYVKA	LAYWET
1	1	0	1.000E+00	0	0
2	0	0	1.000E+00	0	0
3	0	0	1.000E+00	0	0

INTERPRETATION OF LAYER FLAGS:

LAYER	LAYER TYPE (LAYTYP)	INTERBLOCK TRANSMISSIVITY (LAYAVG)	HORIZONTAL ANISOTROPY (CHANI)	DATA IN ARRAY VKA (LAYVKA)	WETTABILITY (LAYWET)
1	CONVERTIBLE	HARMONIC	1.000E+00	VERTICAL K	NON-WETTABLE
2	CONFINED	HARMONIC	1.000E+00	VERTICAL K	NON-WETTABLE
3	CONFINED	HARMONIC	1.000E+00	VERTICAL K	NON-WETTABLE

```

1800 ELEMENTS IN X ARRAY ARE USED BY LPF
18 ELEMENTS IN IX ARRAY ARE USED BY LPF

```

```

SIP5 -- STRONGLY IMPLICIT PROCEDURE SOLUTION PACKAGE
      VERSION 5, 9/1/93 INPUT READ FROM UNIT 19
MAXIMUM OF 50 ITERATIONS ALLOWED FOR CLOSURE
      5 ITERATION PARAMETERS
      2755 ELEMENTS IN X ARRAY ARE USED BY SIP
      150 ELEMENTS IN IX ARRAY ARE USED BY SIP

```

Figure 12. -- GLOBAL file for example problem with parameters—Continued.

```

4555 ELEMENTS OF X ARRAY USED OUT OF      4555
   0 ELEMENTS OF Z ARRAY USED OUT OF      1
168 ELEMENTS OF IX ARRAY USED OUT OF     168
   0 ELEMENTS OF XHS ARRAY USED OUT OF    1

```

SOLUTION BY THE STRONGLY IMPLICIT PROCEDURE

```

-----
MAXIMUM ITERATIONS ALLOWED FOR CLOSURE =      50
      ACCELERATION PARAMETER =      1.0000
HEAD CHANGE CRITERION FOR CLOSURE =    0.10000E-02
SIP HEAD CHANGE PRINTOUT INTERVAL =      1

```

5 ITERATION PARAMETERS CALCULATED FROM SPECIFIED WSEED = 0.00100000 :

```

0.000000E+00 0.822172E+00 0.968377E+00 0.994377E+00 0.999000E+00

```

WETTING CAPABILITY IS NOT ACTIVE IN ANY LAYER

PARAMETERS DEFINED IN THE LPF PACKAGE

```

PARAMETER NAME:HK1          TYPE:HK      CLUSTERS:  1
Parameter value from package file is:  1.00000E-03
      LAYER:  1      MULTIPLIER ARRAY: NONE      ZONE ARRAY: ALL

PARAMETER NAME:HK2          TYPE:HK      CLUSTERS:  1
Parameter value from package file is:  1.00000E-04
      LAYER:  2      MULTIPLIER ARRAY: NONE      ZONE ARRAY: ALL

PARAMETER NAME:HK3          TYPE:HK      CLUSTERS:  1
Parameter value from package file is:  2.00000E-04
      LAYER:  3      MULTIPLIER ARRAY: NONE      ZONE ARRAY: ALL

PARAMETER NAME:VKCB1        TYPE:VKCB    CLUSTERS:  1
Parameter value from package file is:  1.0000
      LAYER:  1      MULTIPLIER ARRAY: MULT1     ZONE ARRAY: ALL

PARAMETER NAME:VKCB2        TYPE:VKCB    CLUSTERS:  1
Parameter value from package file is:  0.50000
      LAYER:  2      MULTIPLIER ARRAY: MULT1     ZONE ARRAY: ALL

```

HYD. COND. ALONG ROWS FOR LAYER 1 WILL BE DEFINED BY PARAMETERS
(PRINT FLAG= 0)

VERTICAL HYD. COND. = 1.00000 FOR LAYER 1

QUASI3D VERT. HYD. COND. FOR LAYER 1 WILL BE DEFINED BY PARAMETERS
(PRINT FLAG= 0)

HYD. COND. ALONG ROWS FOR LAYER 2 WILL BE DEFINED BY PARAMETERS
(PRINT FLAG= 0)

Figure 12. -- GLOBAL file for example problem with parameters—Continued.

VERTICAL HYD. COND. = 1.00000 FOR LAYER 2

QUASI3D VERT. HYD. COND. FOR LAYER 2 WILL BE DEFINED BY PARAMETERS
(PRINT FLAG= 0)

HYD. COND. ALONG ROWS FOR LAYER 3 WILL BE DEFINED BY PARAMETERS
(PRINT FLAG= 0)

VERTICAL HYD. COND. = 1.00000 FOR LAYER 3

1 Well parameters

PARAMETER NAME:WELL1 TYPE:Q
Parameter value from package file is: 1.0000
NUMBER OF ENTRIES: 12

WELL NO.	LAYER	ROW	COL	STRESS FACTOR
1	1	9	8	-5.000
2	1	9	10	-5.000
3	1	9	12	-5.000
4	1	9	14	-5.000
5	1	11	8	-5.000
6	1	11	10	-5.000
7	1	11	12	-5.000
8	1	11	14	-5.000
9	1	13	8	-5.000
10	1	13	10	-5.000
11	1	13	12	-5.000
12	1	13	14	-5.000

1 Drain parameters

PARAMETER NAME:DRN1 TYPE:DRN
Parameter value from package file is: 1.0000
NUMBER OF ENTRIES: 2

DRAIN NO.	LAYER	ROW	COL	DRAIN EL.	STRESS FACTOR
1	1	8	2	0.0000	1.000
2	1	8	3	0.0000	1.000

2 Recharge parameters

PARAMETER NAME:RCH1 TYPE:RCH CLUSTERS: 1
Parameter value from package file is: 3.00000E-08
MULTIPLIER ARRAY: NONE ZONE ARRAY: RCHZONES
ZONE VALUES: 1

Figure 12. -- GLOBAL file for example problem with parameters—Continued.


```
PARAMETER NAME:RCH2          TYPE:RCH    CLUSTERS:  1
Parameter value from package file is:  3.00000E-08
      MULTIPLIER ARRAY: NONE      ZONE ARRAY: RCHZONES
      ZONE VALUES:      2
```

```
  9 PARAMETERS HAVE BEEN DEFINED IN ALL PACKAGES.
(SPACE IS ALLOCATED FOR  500 PARAMETERS.)
```

Figure 12. -- GLOBAL file for example problem with parameters—Continued.

MODFLOW-2000
 U.S. GEOLOGICAL SURVEY MODULAR FINITE-DIFFERENCE GROUND-WATER FLOW MODEL
 VERSION 1.00 06/13/2000

This model run produced both GLOBAL and LIST files. This is the LIST file.

```
#SAMPLE----3 LAYERS, 15 ROWS, 15 COLUMNS; STEADY STATE; CONSTANT HEADS COLUMN 1
#LAYERS 1 AND 2; RECHARGE, WELLS AND DRAINS
  3 LAYERS          15 ROWS          15 COLUMNS
  1 STRESS PERIOD(S) IN SIMULATION
```

```
BAS6 -- BASIC PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 5
      15 ELEMENTS IN IR ARRAY ARE USED BY BAS
```

```
WEL6 -- WELL PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 12
      1 Named Parameters          12 List entries
MAXIMUM OF 15 ACTIVE WELLS AT ONE TIME
      108 ELEMENTS IN RX ARRAY ARE USED BY WEL
```

```
DRN6 -- DRAIN PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 13
      1 Named Parameters          2 List entries
MAXIMUM OF 9 ACTIVE DRAINS AT ONE TIME
      55 ELEMENTS IN RX ARRAY ARE USED BY DRN
```

```
RCH6 -- RECHARGE PACKAGE, VERSION 6, 1/11/2000 INPUT READ FROM UNIT 18
      2 Named Parameters
OPTION 1 -- RECHARGE TO TOP LAYER
      225 ELEMENTS IN RX ARRAY ARE USED BY RCH
      225 ELEMENTS IN IR ARRAY ARE USED BY RCH
```

```
      388 ELEMENTS OF RX ARRAY USED OUT OF          388
      240 ELEMENTS OF IR ARRAY USED OUT OF          240
```

```
1
#SAMPLE----3 LAYERS, 15 ROWS, 15 COLUMNS; STEADY STATE; CONSTANT HEADS COLUMN 1
#LAYERS 1 AND 2; RECHARGE, WELLS AND DRAINS
```

```

          BOUNDARY ARRAY FOR LAYER 1
READING ON UNIT 5 WITH FORMAT: (20I4)

```

	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
1	-1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
2	-1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
3	-1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
4	-1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
5	-1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
6	-1	1	1	1	1	1	1	1	1	1	1	1	1	1	1

Figure 13. -- LIST file for example problem with parameters.

```

7  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
8  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
9  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
10 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
11 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
12 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
13 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
14 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
15 -1  1  1  1  1  1  1  1  1  1  1  1  1  1

```

```

                                BOUNDARY ARRAY FOR LAYER 2
READING ON UNIT 5 WITH FORMAT: (20I4)

      1  2  3  4  5  6  7  8  9 10 11 12 13 14 15
.....
1  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
2  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
3  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
4  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
5  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
6  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
7  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
8  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
9  -1  1  1  1  1  1  1  1  1  1  1  1  1  1
10 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
11 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
12 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
13 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
14 -1  1  1  1  1  1  1  1  1  1  1  1  1  1
15 -1  1  1  1  1  1  1  1  1  1  1  1  1  1

```

BOUNDARY ARRAY = 1 FOR LAYER 3

AQUIFER HEAD WILL BE SET TO 999.99 AT ALL NO-FLOW NODES (IBOUND=0).

INITIAL HEAD = 0.000000 FOR LAYER 1

INITIAL HEAD = 0.000000 FOR LAYER 2

INITIAL HEAD = 0.000000 FOR LAYER 3

OUTPUT CONTROL IS SPECIFIED ONLY AT TIME STEPS FOR WHICH OUTPUT IS DESIRED
HEAD PRINT FORMAT CODE IS 20 DRAWDOWN PRINT FORMAT CODE IS 0
HEADS WILL BE SAVED ON UNIT 0 DRAWDOWNS WILL BE SAVED ON UNIT 0

HYD. COND. ALONG ROWS is defined by the following parameters:
HK1

HYD. COND. ALONG ROWS = 1.000000E-03 FOR LAYER 1

QUASI3D VERT. HYD. COND. is defined by the following parameters:
VKCB1

Figure 13. -- LIST file for example problem with parameters—Continued.

QUASI3D VERT. HYD. COND. = 1.000000E-06 FOR LAYER 1

HYD. COND. ALONG ROWS is defined by the following parameters:
HK2

HYD. COND. ALONG ROWS = 1.000000E-04 FOR LAYER 2

QUASI3D VERT. HYD. COND. is defined by the following parameters:
VKCB2

QUASI3D VERT. HYD. COND. = 5.000000E-07 FOR LAYER 2

HYD. COND. ALONG ROWS is defined by the following parameters:
HK3

HYD. COND. ALONG ROWS = 2.000000E-04 FOR LAYER 3

1
STRESS PERIOD NO. 1, LENGTH = 86400.00

NUMBER OF TIME STEPS = 1

MULTIPLIER FOR DELT = 1.000

INITIAL TIME STEP SIZE = 86400.00

WELL NO.	LAYER	ROW	COL	STRESS RATE
1	3	5	11	-5.000
2	2	4	6	-5.000
3	2	6	12	-5.000

Parameter: WELL1

WELL NO.	LAYER	ROW	COL	STRESS RATE
4	1	9	8	-5.000
5	1	9	10	-5.000
6	1	9	12	-5.000
7	1	9	14	-5.000
8	1	11	8	-5.000
9	1	11	10	-5.000
10	1	11	12	-5.000
11	1	11	14	-5.000
12	1	13	8	-5.000
13	1	13	10	-5.000
14	1	13	12	-5.000
15	1	13	14	-5.000

15 WELLS

DRAIN NO.	LAYER	ROW	COL	DRAIN EL.	CONDUCTANCE
1	1	8	4	10.00	1.000

Figure 13. -- LIST file for example problem with parameters—Continued.

2	1	8	5	20.00	1.000
3	1	8	6	30.00	1.000
4	1	8	7	50.00	1.000
5	1	8	8	70.00	1.000
6	1	8	9	90.00	1.000
7	1	8	10	100.0	1.000

Parameter: DRN1

DRAIN NO.	LAYER	ROW	COL	DRAIN EL.	CONDUCTANCE
8	1	8	2	0.0000	1.000
9	1	8	3	0.0000	1.000

9 DRAINS

RECH array defined by the following parameters:

Parameter: RCH1

Parameter: RCH2

RECHARGE = 3.000000E-08

SOLVING FOR HEAD

31 ITERATIONS FOR TIME STEP 1 IN STRESS PERIOD 1

MAXIMUM HEAD CHANGE FOR EACH ITERATION:

HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL	HEAD CHANGE LAYER,ROW,COL
-22.41 (3, 5, 11)	12.48 (1, 1, 15)	13.39 (3, 1, 14)	48.21 (1, 1, 15)	35.91 (3, 1, 13)
2.482 (1, 9, 14)	1.430 (3, 10, 13)	6.214 (1, 12, 14)	7.411 (3, 11, 14)	13.66 (1, 15, 15)
0.5503 (3, 8, 7)	0.4821 (2, 6, 9)	0.4711 (3, 5, 10)	2.019 (1, 11, 14)	2.302 (3, 5, 13)
0.1108 (1, 13, 12)	0.7058E-01 (3, 12, 11)	0.2819 (1, 14, 14)	0.3141 (3, 13, 14)	0.3320 (1, 15, 15)
0.7853E-02 (1, 13, 12)	0.1586E-01 (2, 11, 11)	0.1777E-01 (3, 11, 10)	0.7910E-01 (1, 14, 14)	0.8500E-01 (3, 7, 14)
0.4169E-02 (1, 13, 14)	0.2555E-02 (3, 14, 15)	0.9769E-02 (1, 14, 14)	0.1082E-01 (3, 13, 14)	0.1030E-01 (1, 15, 15)
0.2429E-03 (1, 13, 12)				

OUTPUT CONTROL FOR STRESS PERIOD 1 TIME STEP 1

PRINT HEAD FOR ALL LAYERS

1

HEAD IN LAYER 1 AT END OF TIME STEP 1 IN STRESS PERIOD 1

Figure 13. -- LIST file for example problem with parameters—Continued.

	1	2	3	4	5	6
	7	8	9	10	11	12
	13	14	15			
1	0.0000	24.94	44.01	59.26	71.82	82.52
	91.91	100.0	106.9	112.6	117.4	121.3
	124.3	126.4	127.4			
2	0.0000	24.45	43.10	57.98	70.17	80.57
	90.12	98.40	105.3	111.0	115.7	119.6
	122.7	124.9	126.1			
3	0.0000	23.45	41.30	55.43	66.78	76.21
	86.51	95.20	102.2	107.6	112.0	116.1
	119.6	122.1	123.4			
4	0.0000	21.92	38.61	51.75	61.79	68.03
	81.34	90.75	97.64	102.5	106.1	110.7
	114.9	117.9	119.4			
5	0.0000	19.73	34.92	47.32	57.69	66.74
	77.09	85.76	92.22	96.15	97.29	103.1
	108.8	112.5	114.3			
6	0.0000	16.51	29.50	40.90	51.30	61.21
	71.19	79.85	86.47	90.82	93.03	94.23
	102.1	106.4	108.4			
7	0.0000	11.55	21.10	31.21	41.40	51.84
	63.08	72.68	79.95	84.92	88.60	91.66
	96.43	99.82	101.8			
8	0.0000	3.483	6.832	16.25	26.30	36.97
	52.59	64.31	72.52	77.25	81.99	85.00
	89.27	91.72	94.33			
9	0.0000	10.54	19.11	28.12	36.92	45.27
	52.95	55.38	65.15	66.07	73.93	73.79
	80.84	80.17	86.49			
10	0.0000	14.62	25.86	35.38	43.49	50.11
	54.93	57.55	62.95	65.55	70.39	72.44
	76.72	78.26	81.79			
11	0.0000	17.11	29.96	40.01	47.78	53.24
	55.81	53.33	60.27	59.29	66.43	65.45
	72.22	71.04	77.62			
12	0.0000	18.68	32.56	43.07	50.81	55.92
	58.33	58.47	61.93	63.18	67.12	68.50
	72.29	73.46	76.85			
13	0.0000	19.67	34.24	45.14	53.01	58.04
	59.91	56.75	62.59	60.91	67.22	65.75
	71.90	70.35	76.48			
14	0.0000	20.27	35.27	46.48	54.61	60.08
	63.17	64.52	67.25	68.79	71.64	73.18
	75.84	77.03	79.09			
15	0.0000	20.56	35.78	47.16	55.48	61.26
	65.02	67.52	69.94	72.01	74.29	76.22
	78.22	79.66	80.82			

1
 HEAD IN LAYER 2 AT END OF TIME STEP 1 IN STRESS PERIOD 1

 1 2 3 4 5 6

Figure 13. -- LIST file for example problem with parameters—Continued.

	7	8	9	10	11	12
	13	14	15			
1	0.0000	24.66	43.73	59.02	71.61	82.32
	91.72	99.86	106.7	112.5	117.2	121.1
	124.1	126.2	127.3			
2	0.0000	24.17	42.83	57.74	69.95	80.36
	89.93	98.22	105.1	110.8	115.5	119.4
	122.6	124.8	125.9			
3	0.0000	23.17	41.03	55.19	66.53	75.77
	86.29	95.02	102.0	107.4	111.8	116.0
	119.5	121.9	123.2			
4	0.0000	21.65	38.34	51.50	61.35	60.17
	80.90	90.55	97.45	102.3	105.4	110.4
	114.8	117.7	119.2			
5	0.0000	19.48	34.65	47.07	57.44	66.30
	76.85	85.57	92.00	95.41	91.09	102.1
	108.6	112.4	114.2			
6	0.0000	16.27	29.24	40.65	51.07	60.98
	70.98	79.65	86.28	90.54	92.06	86.23
	101.7	106.2	108.3			
7	0.0000	11.38	20.95	31.05	41.25	51.70
	62.90	72.48	79.76	84.73	88.35	91.24
	96.22	99.65	101.6			
8	0.0000	4.209	8.330	17.58	27.58	38.25
	52.94	64.19	72.34	77.12	81.81	84.86
	89.10	91.59	94.17			
9	0.0000	10.38	18.96	27.98	36.79	45.16
	52.86	56.13	65.08	66.79	73.87	74.48
	80.77	80.84	86.38			
10	0.0000	14.40	25.61	35.15	43.27	49.91
	54.76	57.48	62.79	65.49	70.24	72.37
	76.57	78.20	81.64			
11	0.0000	16.87	29.70	39.78	47.56	53.05
	55.68	54.09	60.20	60.04	66.37	66.18
	72.16	71.75	77.51			
12	0.0000	18.43	32.31	42.85	50.60	55.73
	58.16	58.41	61.78	63.12	66.98	68.44
	72.15	73.40	76.69			
13	0.0000	19.42	33.98	44.91	52.80	57.85
	59.78	57.50	62.53	61.65	67.16	66.48
	71.84	71.06	76.37			
14	0.0000	20.02	35.02	46.26	54.41	59.88
	62.99	64.39	67.08	68.66	71.48	73.06
	75.68	76.91	78.93			
15	0.0000	20.30	35.52	46.94	55.28	61.07
	64.84	67.34	69.76	71.84	74.11	76.04
	78.04	79.49	80.65			

1

HEAD IN LAYER 3 AT END OF TIME STEP 1 IN STRESS PERIOD 1

1	2	3	4	5	6

Figure 13. -- LIST file for example problem with parameters—Continued.

	7	8	9	10	11	12
	13	14	15			
1	1.800	24.34	43.36	58.70	71.33	82.06
	91.48	99.63	106.5	112.3	117.0	120.9
	123.9	126.0	127.1			
2	1.764	23.85	42.46	57.42	69.66	80.07
	89.68	97.99	104.9	110.6	115.3	119.2
	122.4	124.6	125.7			
3	1.691	22.86	40.67	54.87	66.20	75.28
	85.98	94.77	101.7	107.2	111.5	115.7
	119.3	121.7	123.0			
4	1.578	21.35	37.98	51.17	60.85	62.69
	80.41	90.28	97.19	101.9	104.1	110.0
	114.5	117.5	119.0			
5	1.415	19.18	34.30	46.75	57.10	65.80
	76.54	85.30	91.67	94.17	77.46	100.7
	108.2	112.1	114.0			
6	1.176	15.99	28.91	40.33	50.76	60.67
	70.70	79.38	86.01	90.12	90.60	88.55
	101.2	106.0	108.0			
7	0.8273	11.21	20.79	30.88	41.09	51.55
	62.67	72.22	79.50	84.46	87.98	90.77
	95.94	99.41	101.4			
8	0.4331	5.131	10.19	19.27	29.19	39.84
	53.40	64.07	72.11	76.95	81.58	84.68
	88.88	91.44	93.95			
9	0.7543	10.22	18.82	27.84	36.66	45.06
	52.78	57.03	65.02	67.64	73.81	75.31
	80.72	81.64	86.24			
10	1.039	14.13	25.29	34.85	42.99	49.65
	54.54	57.44	62.61	65.44	70.05	72.33
	76.39	78.15	81.43			
11	1.224	16.59	29.37	39.47	47.28	52.79
	55.53	55.01	60.16	60.94	66.33	67.06
	72.13	72.60	77.38			
12	1.341	18.15	31.97	42.54	50.32	55.47
	57.94	58.37	61.60	63.08	66.80	68.41
	71.97	73.36	76.49			
13	1.415	19.14	33.65	44.61	52.53	57.60
	59.63	58.39	62.48	62.54	67.12	67.35
	71.80	71.90	76.24			
14	1.460	19.73	34.68	45.96	54.13	59.63
	62.76	64.24	66.87	68.52	71.27	72.91
	75.47	76.77	78.71			
15	1.481	20.01	35.18	46.63	55.00	60.81
	64.59	67.11	69.52	71.61	73.87	75.82
	77.81	79.27	80.42			

1

Figure 13. -- LIST file for example problem with parameters—Continued.

VOLUMETRIC BUDGET FOR ENTIRE MODEL AT END OF TIME STEP 1 IN STRESS PERIOD 1

CUMULATIVE VOLUMES	L**3	RATES FOR THIS TIME STEP	L**3/T
IN:		IN:	
STORAGE =	0.0000	STORAGE =	0.0000
CONSTANT HEAD =	0.0000	CONSTANT HEAD =	0.0000
WELLS =	0.0000	WELLS =	0.0000
DRAINS =	0.0000	DRAINS =	0.0000
RECHARGE =	13608000.0000	RECHARGE =	157.5000
TOTAL IN =	13608000.0000	TOTAL IN =	157.5000
OUT:		OUT:	
STORAGE =	0.0000	STORAGE =	0.0000
CONSTANT HEAD =	4326522.0000	CONSTANT HEAD =	50.0755
WELLS =	6480000.0000	WELLS =	75.0000
DRAINS =	2801081.2500	DRAINS =	32.4199
RECHARGE =	0.0000	RECHARGE =	0.0000
TOTAL OUT =	13607603.0000	TOTAL OUT =	157.4954
IN - OUT =	397.0000	IN - OUT =	4.5929E-03
PERCENT DISCREPANCY =	0.00	PERCENT DISCREPANCY =	0.00

TIME SUMMARY AT END OF TIME STEP 1 IN STRESS PERIOD 1					
	SECONDS	MINUTES	HOURS	DAYS	YEARS
TIME STEP LENGTH	86400.	1440.0	24.000	1.0000	2.73785E-03
STRESS PERIOD TIME	86400.	1440.0	24.000	1.0000	2.73785E-03
TOTAL TIME	86400.	1440.0	24.000	1.0000	2.73785E-03

1

Figure 13. -- LIST file for example problem with parameters—Continued.

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