

PHREEQCI—A Graphical User Interface for the Geochemical Computer Program PHREEQC

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by Scott R. Charlton, Clifford L. Macklin, and David L. Parkhurst

Abstract

PHREEQCI is a Windows-based graphical user interface for the geochemical computer program PHREEQC. PHREEQCI provides the capability to generate and edit input data files, run simulations, and view text files containing simulation results, all within the framework of a single interface. PHREEQC is a multipurpose geochemical program that can perform speciation, inverse, reaction-path, and 1D advective reaction-transport modeling. Interactive access to all of the capabilities of PHREEQC is available with PHREEQCI. The interface is written in Visual Basic and will run on personal computers under the Windows(3.1), Windows95, and WindowsNT operating systems.

INTRODUCTION

PHREEQC (Parkhurst, 1995) is a computer program that can perform several types of geochemical modeling including speciation, inverse, reaction-path, and 1D advective reaction-transport modeling. PHREEQCI is a Windows-based interface that provides access to all the capabilities of PHREEQC. Data for PHREEQC are entered through a series of keyword data blocks; each data block contains a specific type of information. For example, the **SOLUTION** keyword data block contains all the information to define the composition of an aqueous solution; **EQUILIBRIUM_PHASES** keyword data block defines the identity and amount of each mineral in a phase assemblage. The keyword data blocks are used to define the data needed for each type of modeling. Additional keyword data blocks are used to define output options and to define and modify the chemical database. The following sections of the Introduction give an overview of the use of the keyword data blocks for PHREEQC.

Speciation Modeling

Speciation modeling uses a chemical analysis of a water to calculate the distribution of aqueous species using an ion-association aqueous model. The most important results of speciation calculations are saturation indices for minerals, which indicate the saturation state of each mineral relative to the water. Normally, speciation modeling requires only a **SOLUTION** keyword data block for each water analysis for which saturation indices are to be calculated.

Inverse Modeling

Inverse modeling is used to deduce geochemical reactions that account for the change in chemical composition of water along a flow path. At least two chemical analyses of water are needed at different points along the flow path as well as a set of phases that are potentially reactive along the flow path. From the analyses and phases, mole balance models are calculated. A mole balance model is a set of mole transfers of phases that accounts for the change in composition along the flow path. Normally, only **SOLUTION** keyword data blocks and an **INVERSE_MODELING** keyword data block are needed for inverse modeling calculations. However, additional reactant phases may need to be defined with **PHASES** or **EXCHANGE_SPECIES** keyword data blocks.

Reaction-Path Modeling

Reaction-path modeling can be used to simulate a wide range of geochemical reactions. In general, these reactions can be divided into equilibrium (or reversible) reactions and irreversible reactions. Reversible reactions include equilibration of a solution (**SOLUTION** keyword) with one or more of the following entities: a phase assemblage (**EQUILIBRIUM_PHASES** keyword), a set of ion

exchangers (**EXCHANGE** keyword), a set of surface complexers (**SURFACE** keyword), and (or) a fixed-pressure gas phase (**GAS_PHASE** keyword).

Irreversible reactions include mixing of solutions (**MIX** keyword), addition (or removal) of fixed amounts of specified reactants (**REACTION** keyword), and changing temperature (**REACTION_TEMPERATURE** keyword).

The same keyword may be used several times to define reactants; an integer number is used to identify each keyword data block. Thus, the user can define three different solutions and assign, for example, the integers 1, 10, and 20 to identify them—**SOLUTION** 1, **SOLUTION** 10, and **SOLUTION** 20. Integer identifiers also are used with **EQUILIBRIUM_PHASES**, **EXCHANGE**, **SURFACE**, **GAS_PHASE**, **MIX**, **REACTION**, and **REACTION_TEMPERATURE** data blocks.

Reaction-path modeling is used to simulate reactions that can be thought of as occurring in a beaker. A solution, or mixture of solutions, is placed in the beaker along with any other equilibrium reactants (phase assemblage, exchange assemblage, surface assemblage, or gas phase), irreversible reactions are applied (**REACTION** and **REACTION_TEMPERATURE**), and the new equilibrium state is calculated.

A key concept for PHREEQC reaction-path calculations is a “simulation.” A simulation is defined to be a set of keyword data blocks terminated by the **END** keyword. Reversible and irreversible reactants are defined within a simulation either by keyword data blocks (first occurrence within a simulation of **SOLUTION**, **EQUILIBRIUM_PHASES**, **REACTION**, and other types of keyword data blocks) or by the **USE** keyword, which specifies the use of a reactant that has been defined in a previous simulation. New compositions of the solution, phase assemblage, exchange assemblage, surface assemblage, and gas phase that are calculated by a reaction simulation can be saved for use in subsequent simulations with the **SAVE** keyword.

Reaction-path modeling may include the use of any of the following keyword data blocks: **SOLUTION**, **EQUILIBRIUM_PHASES**, **EXCHANGE**, **SURFACE**, **GAS_PHASE**, **MIX**, **REACTION**, **REACTION_TEMPERATURE**, **SAVE**, and **USE**.

1D Advective Reaction-Transport Modeling

Reaction-transport modeling simulates advection and chemical reactions as water moves through a one dimensional column. The column is divided into a number of cells, n , which is defined by the user. The cells are numbered 1 through n , and these cells initially contain solutions with identifying numbers 1 through n . A solution composition for each of these integers must have been defined by **SOLUTION** data blocks (note that data for a range of numbers can be defined simultaneously, for example **SOLUTION** 1-15) or the **SAVE** keyword (for example, **SAVE solution** 1-15). The cells may also contain other reversible or irreversible reactants. For a given cell number, i , if a phase assemblage, exchange assemblage, surface assemblage, gas phase, mixture, reaction, or reaction temperature with identifying number i has been defined, then it is automatically present in cell i during the transport calculation. Thus, the initial conditions and the set of reactants of each cell can be defined individually, which allows the flexibility to simulate a variety of chemical conditions throughout the column.

Advection is simulated by moving the solution in each cell to the next higher numbered cell; **SOLUTION** 0 must be defined and it is moved into cell 1; the solution from cell 1 is moved to cell 2, and so on, until the solution from cell $n-1$ is moved to cell n . At this point, the solution in cell n is discarded. All reversible and irreversible reactants except the solution remain in their original cells. After each cell has received its new solution, irreversible reactions are applied within the cell, the solution is equilibrated with the reversible reactants, and the equilibrium compositions of the solution and reversible reactants are saved.

Advective reaction-transport modeling uses the **TRANSPORT** keyword data block. Usually, one or more simulations precede the **TRANSPORT** simulation. These preceding simulations often use many of the keywords described in reaction-path modeling to define the initial solutions and reactants of the column and to define the infilling solution (**SOLUTION** 0).

Printing Options

A large amount of information is produced by speciation, reaction-path, and reaction-transport calculations. By default, all results are written to a single output file, which may become very large, especially

for reaction-transport calculations. The data written to the output file can be limited by use of the **PRINT** keyword data block and by options within the **TRANSPORT** keyword data block. In addition, a “selected-output file” that contains selected results can be specified. This file contains columns of data for user-specified items. For example, the total concentration of iron and the saturation index of calcite could be specified to be written to the selected-output file for each cell and time step of a transport calculation. The name of the file and the data items to be written are defined with the **SELECTED_OUTPUT** keyword data block. The selected-output file consists of columns of data and can be easily imported into a spreadsheet program for analysis and graphing.

Database Files: Elements, Aqueous Species, Exchange Species, Surface Species, and Pure Phases

PHREEQC uses a database file to define the set of elements and element redox states, ion exchangers, surface complexers, and pure phases that are used in calculations. The database file contains all of the stoichiometry and thermodynamic data for aqueous species, exchange species, surface species, and pure phases. These data are defined through the following keyword data blocks:

SOLUTION_MASTER_SPECIES,
SOLUTION_SPECIES,
EXCHANGE_MASTER_SPECIES,
EXCHANGE_SPECIES,
SURFACE_MASTER_SPECIES,
SURFACE_SPECIES, and **PHASES**. Custom database files can be written using these keywords. In addition, database files can be modified at run time through the use of these keywords in the input file.

Three database files are distributed with PHREEQC and PHREEQCI, *phreeqc.dat*, *wateq4f.dat*, and *minteq.dat*. The file *phreeqc.dat* is derived from the database file for the geochemical program PHREEQE (Parkhurst and others, 1980), but the data are consistent with *wateq4f.dat* for the minerals and species that are common to both. The file *phreeqc.dat* contains data for the elements Al, B, Ba, Br, C, Ca, Cd, Cl, Cu, F, Fe, K, Li, Mg, Mn, N, Na, P, Pb, S, Sr, Si, and Zn. The file *wateq4f.dat* is derived from the database file for WATEQ4F (Ball and Nordstrom, 1991) and contains all of the elements from *phreeqc.dat* plus As, Cs, I, Ni, Rb,

Se, and U. In addition, *wateq4f.dat* has data for organic ligands labeled Fulvate and Humate. The file *minteq.dat* is translated from the data files of MINTEQA2 (Allison and others, 1990).

THE INTERFACE

When PHREEQCI is installed on a computer, a new icon is created in the Windows environment (see How to Obtain and Install PHREEQCI below). The interface can be invoked by double clicking on the icon. PHREEQCI is installed in a directory of the user's choice. For this manual, it is assumed that the program is installed in the directory *c:\phreeqci*. The program that is executed by double clicking the icon is *c:\phreeqci\phreeqci.exe*. After invoking the program, the main screen for PHREEQCI appears (fig. 1).

The main screen for PHREEQCI has a button for each keyword (fig. 1) that produces an input screen for the keyword data block. After data have been entered in the screen for a keyword data block, the data are stored in a temporary copy of the input data file and the main screen reappears. **END** keywords can be added with the “End of Simulation” button that spans the main screen at the bottom (fig. 1). Individual keyword data blocks may be retrieved and edited through the “Field Editor” button on the main screen (fig. 1). The entire input data set may be edited with a user-specified editor (see Options below) through the “Line Editor” button (fig. 1). Once all of the keywords for all the simulations are entered, PHREEQC can be run with the “Run Current File” button. The top border of the main screen lists the name of the current input file and lists the number of simulations in the file (see Reaction-Path Modeling above).

Any item on a screen that has an underlined letter can be invoked through the keyboard by pressing the “Alt” key and the letter key simultaneously.

Pull-Down Menus on the Main Screen

Four pull-down menus are located at the top of the main screen: “File”, “New Keyword”, “Options”, and “Help” (fig. 1).

File

The “File” pull-down menu on the main screen (fig. 1) allows the selection of a new (“New”) or exist-

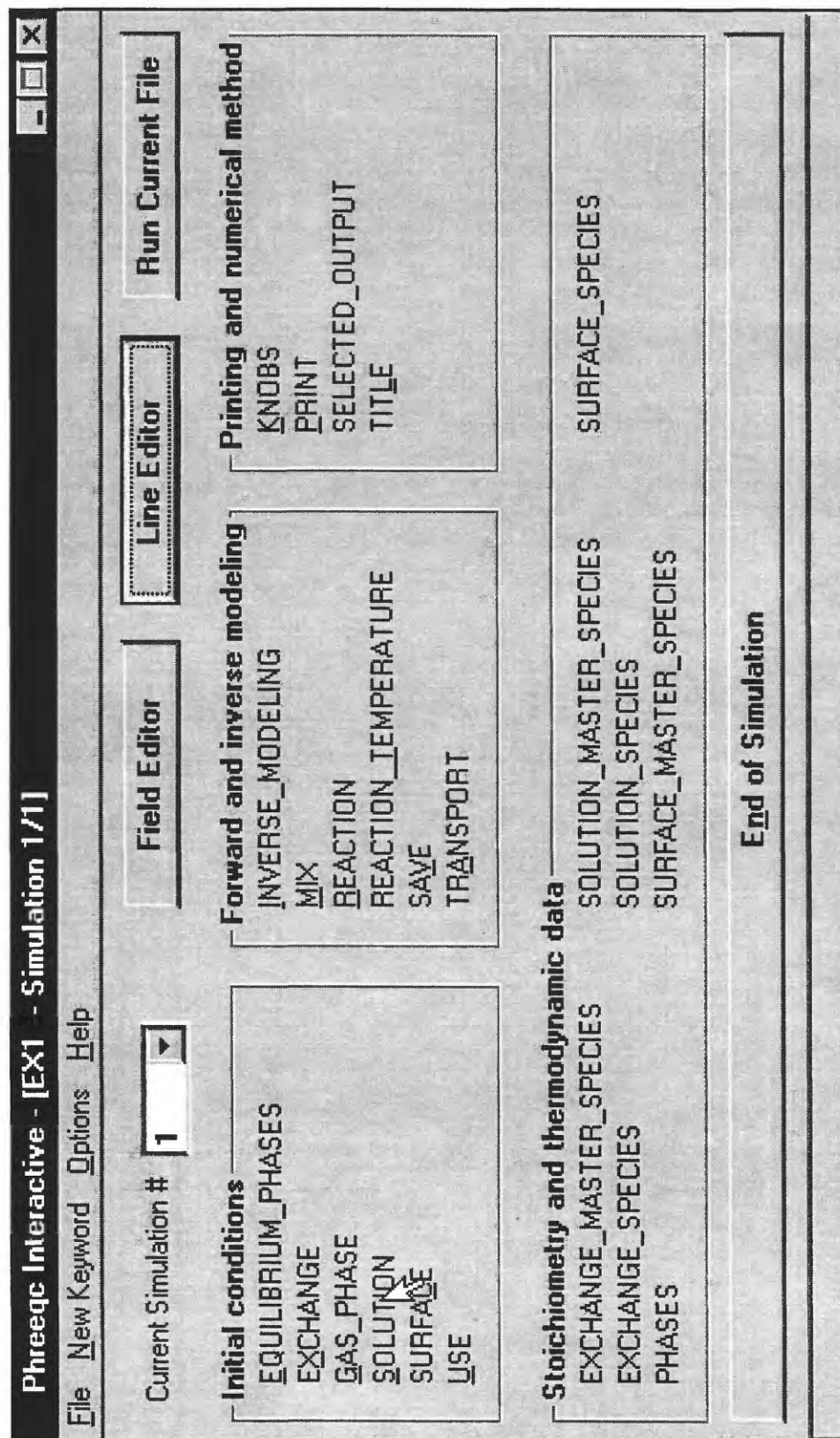


Figure 1.—Main screen of PHREEQCI.

ing ("Open") file as the input data file. It also allows the current input data file to be saved ("Save") or renamed ("Save As"). The "Exit" item in the menu closes the program PHREEQCI and returns to Windows.

New Keyword

The "New Keyword" pull-down menu on the main screen (fig. 1) allows a selection of any keyword screen using only keystrokes—that is, without the use of the mouse.

Options

The "Options" pull-down menu is used to set preferences for a browser, text editors, and default database file.

The browser is used by the help facility to view the help files. Commonly, Netscape, Internet Explorer, or Mosaic are available to use as browsers. If no browser is available, then the help facility will not function, but the rest of the interface will function normally. Two boxes are used to define the browser in the "Options" window: (1) The box labeled "HTML Browser" is used to define the browser; on some systems it may be necessary to give the entire path to the browser program, for example *c:/mosaic/mosaic.exe*, and (2) The box labeled "HTML Browser Options" allows any special options needed to invoke the browser to be defined; for example to invoke Mosaic in stand-alone mode, "-s" should be entered in this box.

The user's preference for text editors for input and output files can be specified in entry boxes within the box labeled "Text Editors" of the "Options" window. The box labeled "Input" is used to define the text editor that is used to edit the input file. Two radio buttons allow the selection of the default Visual Basic editor ("Built-in") or another editor ("User Defined") that is available on the computer. If "User Defined" is selected, the name, or possibly the pathname to the executable file, of the editor, must be entered. A browser is available to locate the pathname to the executable file if the desired editor can not be located through the default "PATH". The box labeled "Output" allows the selection of an editor to use with output files. The default text editor is Notepad. With the Windows(3.1) and Windows95 operating systems, this text editor allows only a limited-size file to be opened and cannot handle long output files. Selection of a more versatile text editor is desirable. As with the user-defined editor

for the input file, a browser is available to locate the pathname to the executable file if the desired editor can not be located through the default PATH.

Finally, the chemical database file for calculations is selected through the "Default Database File" box of the "Options" window. The three database files distributed with PHREEQCI, *phreeqc.dat*, *wateq4f.dat*, and *minteq.dat*, are stored in the top level directory of the PHREEQCI installation and may be used by typing the selected name in the entry box. If the user has developed a custom database file, it can be selected with the "browse" button next to the entry box.

Help

The "Help" pull-down menus on all screens use the HTML browser (see Options above) to read on-line documentation. The "Help" pull-down menu on the main screen (fig. 1) has three items: "PHREEQCI" displays the first page of this manual; "User's Guide to PHREEQC" displays the first page of the user's manual for PHREEQC (Parkhurst, 1995); and "About PHREEQC Interactive" displays the version number of PHREEQCI.

Keyword Screens

Each keyword of PHREEQC is listed on the main screen of PHREEQCI. Clicking on a keyword, or if the keyword has an underlined letter, pressing the Alt key and the letter key simultaneously will produce a new screen for entry of that keyword data block. The keyword data block will be inserted into the simulation number given in the "Current Simulation #" box shown in the upper left of the main screen. Clicking "End of Simulation" places an **END** keyword in the input file and advances the current simulation number, but does not produce a new screen.

An example keyword screen for the **SOLUTION** keyword is shown in figure 2. The "Help" menu at the top of a keyword screen (fig. 2) invokes the HTML browser (see Options above) and displays the description of data input for the keyword from the PHREEQC manual (Parkhurst, 1995).

All keyword screens have the three buttons "Done", "Cancel", and "Clear All" in the upper right-hand corner (fig. 2). The "Done" button is used when all data have been entered for the keyword data block. After the button is pushed, the data are written to a temporary copy of the input file and control is returned to

SOLUTION - [EX1 -- Simulation 1/2] X

Help

SOLUTION

Number through Description Done
Cancel
Clear All

pH Equilibrium Sat. index ☐ Charge balance

pe Equilibrium Sat. index

Temperature

Redox Density

Units

Individual element input

Element/State	Units	Redox	Concentration	As	GFw	Phase equilibrium	Sat. index	Charge balance
<input type="text" value="O(0)"/>	<input type="text"/>	<input type="text"/>	<input type="text" value="1"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="checkbox"/>
<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="checkbox"/>
<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="checkbox"/>
<input type="text" value="O2(g)"/>	<input type="text"/>	<input type="text"/>	<input type="text" value="-0.7"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="checkbox"/>

List of elements

S(6)	2712							
N(5)	0.29	as NO3						
N(-3)	0.03	as NH4						
U	3.3	ppb						
O(0)	1	O2(g)	-0.7					

Delete Selected Entry

Figure 2.—Keyword screen for **SOLUTION** keyword data block.

the main screen. The “Cancel” button removes the keyword screen and no data are stored to the input file. The “Clear All” button deletes all information in all boxes of the keyword screen and reinitializes any default values.

Entry boxes can be selected by a mouse click, a tab, or a “shift tab”. The tab key cycles the cursor through the entry boxes in sequential order. The shift and tab keys pressed simultaneously move the cursor to entry boxes in reverse sequential order.

Data for any entry boxes can be entered directly from the keyboard after being selected by a mouse click, a tab, or a shift tab. Many entry boxes have adjoining list boxes, which are buttons with downward-pointing arrows. A mouse click on a list box or pressing the Alt and the down arrow key simultaneously while the entry box is selected will display a list of possible entries for the adjoining entry box. Pointing the mouse cursor to an entry and clicking puts the selected entry in the entry box. Alternatively, while the list of possible entries is displayed, pressing a sequence of letters on the keyboard will advance the selection to an entry that begins with that sequence of letters; pressing the enter key will put the selected entry in the entry box.

Many keyword screens define lists of information. For example, in the **SOLUTION** keyword screen, a list of elements and concentrations must be entered (fig. 2). The list of the **SOLUTION** screen will be described here, but all lists are entered and edited with the same logic. The data for one element are entered in the “Individual element input” boxes (for example the “Element/State” and “Concentration” boxes in fig. 2). The “Enter” button is clicked to add the new entry to the “List of elements” (fig. 2). Any entry in the “List of elements” may be deleted by selecting it with a single mouse click followed by a click on the “Delete Selected Entry” button (fig. 2). Any entry from the “List of elements” may be selected and edited by double clicking with the mouse. After selecting an item from the “List of elements” by double clicking, the entry boxes of “Individual element input” (“Element/State”, “Concentration”, “Units”, and others, fig. 2) are populated with the selected item and the data in the entry boxes can be edited. Another click of the “Enter” button replaces the selected item in the “List of elements” with the edited data.

Field Editor

Any keyword data block that is in the input data file can be edited with the appropriate keyword screen by using the “Field Editor” button on the main screen. A new screen with two entry boxes appears. First, the correct simulation must be selected with the “Simulations” entry box, and then the desired keyword can be selected with the “Keyword” entry box.

Line Editor

The entire input file can be edited at any time with a user-defined editor by using the “Line Editor” button on the main screen. (The editor for the “Line Editor” button is specified in the Options menu, see Options above.) After exiting the line editor, control returns to the main screen.

Input to PHREEQC is free format and fields are delimited by spaces or tabs. In general, the order of keywords within a simulation and, in most cases, the order of lines within a keyword are irrelevant, so the input file can be edited relatively easily. However, it is possible to introduce input errors that cause the data blocks not to be interpretable by PHREEQC and PHREEQCI. In this case, the user has to correct the errors in the line editor, or remove the erroneous keyword data block with the line editor and reenter the data through the keyword screen.

The line editor is the only way to delete an entire keyword data block, to add an **END** keyword between previously defined keywords, or to move keyword data blocks between simulations.

After familiarization with PHREEQCI, the line editor can be used for simple modifications of an input file or even for much of the data entry.

Run Current File

Once all of the keywords for all the simulations are entered, PHREEQC can be run with the “Run Current File” button. At this point, a window labeled “Save As” appears and is used to select the directory and name of the file in which to store all the keyword data that have been entered. Next a window labeled “Save Output As” appears and is used to select the directory and name of the output file that is generated by PHREEQC. PHREEQC is then run and the output file is displayed to the screen

in the text editor specified with the Options menu (see Options above). After viewing results, exiting from the text editor returns control to the main screen of PHREEQCI.

FILES

The user has control over four files that are used or produced by a PHREEQCI run: (1) input file, (2) output file, (3) selected-output file, and (4) database file.

Input File

The input file contains a set of keywords and associated data that define the set of geochemical calculations to be done. An input file is required for all calculations. The name of the input file is selected either with the "File" menu on the main screen of the program, or at the time the "Run Current File" button on the main screen is selected. All information that has been read from an existing input file or input through keyword screens or the line editor is written to the input file. The "File" menu on the main screen allows saving, changing, and renaming the input file.

Output File

The output file contains all results of PHREEQC calculations and is required for all calculations. The name of the file is selected after the "Run Current File" button on the main screen is selected. By default, the output file name is the input file name with ".out" appended. The results that are written to the output file can be changed by options in the **PRINT** and **TRANSPORT** keyword data blocks.

Selected-Output File

A file suitable for importing into a spreadsheet program can be defined through the **SELECTED_OUTPUT** keyword data block. The name of the file can be defined (default is *selected.out*) and the data items to be included in the file can be selected. When **SELECTED_OUTPUT** keyword data block is used, a set of data items, including the numbers of reactants, pH, temperature, ionic strength, and other items, is printed to the selected-output file. In addition, the following items can be written to the selected-out-

put file for every PHREEQC calculation: (1) the total molality of any element or element redox state, (2) the molality of any aqueous, exchange, or surface species, (3) the log of the activity of any aqueous, exchange, or surface species, (4) the amount and mole transfer of any mineral in the reaction phase assemblage, (5) the saturation index for any mineral, and (6) the amount of any gas in the gas phase.

Database File

The definitions of the elements and element redox states, ion exchangers, and surface complexers are normally included in the database file. In addition, the stoichiometry and thermodynamic data for aqueous, exchange, and surface species and for pure phases are normally defined in the database file.

Three database files are distributed with PHREEQCI, *phreeqc.dat*, *wateq4f.dat*, and *minteq.dat* (see Database Files: Elements, Aqueous Species, Exchange Species, Surface Species, and Pure Phases above). One of these database files or a custom database file can be selected through the Options menu on the main screen.

HOW TO OBTAIN AND INSTALL PHREEQCI

The latest version of the software described in this report and a Postscript file of this manual can be obtained from the web site <http://water.usgs.gov/software/>. Many other hydrologic computer programs are also available at this site.

The latest version of PHREEQCI may also be obtained by anonymous ftp from the Internet address: [brrcrftp.cr.usgs.gov](ftp://brrcrftp.cr.usgs.gov) (136.177.112.5). The files reside in the directory */geochem/pc/phreeqci*. A typical anonymous ftp session follows:

```
% ftp brrcrftp.cr.usgs.gov
Name: anonymous
Password: e-mail address (replaced with e-mail address)
ftp> cd geochem/pc/phreeqci (change directory)
ftp> ls (list files in directory)
pcix-xx.exe ("x-xx" represents the version number)
ftp> type binary (eliminate translation for binary files)
ftp> get pcix-xx.exe (transfer the file)
ftp> quit (quit ftp)
```

Alternatively, the documentation and current version of the software can be ordered from the following address:

U.S. Geological Survey
NWIS Program Office
437 National Center
Reston, VA 22192
(703) 648-5695

Additional copies of this report are available from:

U.S. Geological Survey
Branch of Information Services
Box 25286
Denver, CO 80225-0286

For additional information, write to the address on page *ii* of this report.

Downloading and Installation

Two methods for downloading and installing PHREEQCI are available. The first method downloads a single self-extracting file, which is extracted in a temporary directory on the hard disk, and PHREEQCI is then installed from this temporary directory. The second method downloads three self-extracting files, which are then extracted onto three 1.44MB floppy diskettes, and PHREEQCI is installed from the floppy diskettes.

Directory Installation

1. Create a temporary directory on the hard drive.
2. Download *pcix-xx.exe* into the temporary directory, where *x-xx* is the version number. (Note: Be sure to use binary transfer if using FTP.)
3. Extract the installation files into the temporary directory by executing *pcix-xx.exe*.
4. Run *setup.exe*.
5. Remove the temporary directory from the hard drive.

Floppy Diskette Installation

1. Create a temporary directory on the hard drive.
2. Download *disk1.exe*, *disk2.exe*, and *disk3.exe*. (Note: Be sure to use binary mode if using FTP.)

3. Create the installation diskettes by extracting the three files onto three blank formatted (1.44MB) floppy diskettes.

For example:

c:\tmp\disk1 a:
c:\tmp\disk2 a:
c:\tmp\disk3 a:

4. Run *setup.exe* from the first floppy diskette.
5. Remove the temporary directory from the hard drive.

WIN32S

PHREEQCI contains a 32-bit component that runs directly with Windows95 and WindowsNT operating systems. However, if the operating system is Windows(3.1), *win32s* is required and *win32s* must be installed before installing PHREEQCI. *Win32s* is available from the server brrcrftp.cr.usgs.gov in the directory */geochem/pc/phreeqi/win32s*. See the *readme.txt* file in that directory for downloading and installation instructions. For more information on *win32s*, please see the following URL: <http://www.ncsa.uiuc.edu/SDG/Software/mosaic-w/faq/win32.html>.

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