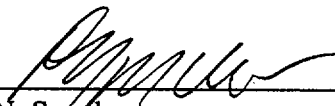


DESIGN DOCUMENT (DD)
for
TOUGHREACT Version 2.2

SAN: LBNL-1999-141
Document Identifier: 10154-DD-2.2-01
STN: 10154-2.2-00
SMN: 10154-MED-2.2-00


Prepared by:


N. Spycher
Software Developer

Date 5/1/00

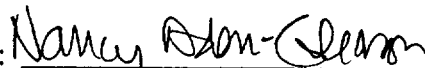
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Verified by:


S. Mukhopadhyay
Technical Reviewer


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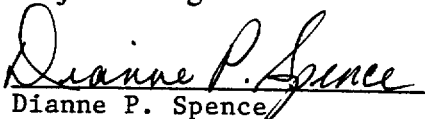

N. Aden-Gleason
Management Reviewer

Date 5/2/00

Approved by:


G. S. Bodvarsson
Project Manager

Date 5/3/00


Dianne P. Spence
ITSMA

Date 5/12/00

**PRELIMINARY DRAFT
INFORMATION ONLY**

CHANGE HISTORY

Revision Number	Effective Date	Description of Change
00	03/12/00	Initial issue of CP1 documents prior to ITSMA review.
01	04/21/00	Initial issue of CP1 documents following ITSMA review. The changes were the addition of the "QA" designator, this Change History, extensive changes to appendices to include referenced material and the resulting change in page numbers.

The design of TOUGHREACT V2.2 has been developed to meet the requirements of the software project's Software Activity Plan (SAP) 10154-SAP-2.2-01 and Requirements Document (RD) 10154-RD-2.2-01 for TOUGHREACT V2.2 as well as LBNL Master Planning Document (MPD) YMP-MPD-UZ-1.0, *UZ Flow and Transport Modeling FY99*. In accordance with AP-SI.1Q, Rev. 2, ICN 4, *Software Management*, this Design Document (DD) is followed by Installation Test Plan (ITP) 10154-ITP-2.2-01 for TOUGHREACT V2.2.

I. PURPOSE

TOUGHREACT V2.2 is an upgrade of TOUGHREACT V2.1 (STN: 10042-2.1-01) (previously baselined and qualified, Spycher et al. 1999). In TOUGHREACT V2.2, the following modifications and additions are to be made (see Requirements Document 10154-RD-2.2-01) as listed below:

1. All upgrades necessary for consistency with TOUGH2 EOS3 v1.4 (STN: 10007-1.4-01) (Wu et al. 1999): active-fracture model, downstream weighting option, capillary pressure linear extrapolation at low liquid saturations, changes in relative permeability based on a modified Brooks-Corey model;
2. Restart option for reactive transport;
3. New option for calculating gas diffusion coefficients as function of temperature and pressure;
4. Use input tortuosity to weight molecular diffusion coefficients;
5. New options for calculating effective surface areas;
6. New option for precipitation rate law;
7. Time dependent thermal conductivity factors;
8. New conjugate gradient stabilized solver;
9. Carmen-Kozeny porosity-permeability coupling option;
10. Modified output options and formats (separate plot output file for gases, option to output gas and mineral amounts in volume fraction, output additional variables such as porosity and permeability in plot files).

II. IMPLEMENTATION STANDARDS AND ENVIRONMENT SPECIFICATIONS

II.1 Hardware Platform(s) and Operating System(s)

This software is intended to be run on Unix workstations and servers. See Appendix B. The hardware system needs to be, at a minimum, a 200 MHz CPU with 64 MB RAM and 1 GB disk.

II.2 Installation and Validation Methodology

The installation and validation methodology shall consist of sample problems for installation checkout and specified test cases that demonstrate that the new features of the software meet the requirements. For installation, there shall be two sample problems with input and output listed in the Validation Test Report for this software. For validation, there shall be several test cases as listed in the Validation Test Plan for this software.

II.3 Programming Standards

The code shall be written in FORTRAN77.

III. SOFTWARE STRUCTURE

The TOUGHREACT V2.2 software structure is defined in Appendix C. The discussion below primarily relates to the upgrades to be incorporated into version 2.2 of TOUGHREACT.

III.1 Software Processes

The software processes to be incorporated into TOUGHREACT V2.2 are derived from testable requirements in the RD, Section I.1. They include the ten upgrades listed above in Section I.

III.2 Process Functions and Software Modules

See description of the process functions and modules in Appendices C, D, and F.

III.3 Internal Interfaces, Control Logic, Data Structure, and Data Flow

These issues are described in Appendix C.

III.4 Hierarchical Structure, Design Entities, and Design Dependencies

These issues are described in Appendix C.

IV. SYSTEM INPUTS AND OUTPUTS

These will be essentially the same as for the previous version 2.1, described in Appendices D and E with some input and output modifications to account for the new options listed in Section I, described in Appendix F.

IV.1 Electronic, Manual, and Printed Inputs

The software shall read (electronic) input files as described in Appendix D, Sections D.6.1 and D.6.2. Modifications for TOUGHREACT V2.2 are described in Appendix F.

IV.2 Input Name, Functional Description, and Source

The input name, functional description, and source of the input files are described in Appendix D, Sections D.6.1.1 and D.6.2. Modifications for TOUGHREACT V2.2 are described in Appendix F.

IV.3 Input Frequency and Format

The frequency of input shall be once per run. Runs may be restarted after reaching a stopping point. The formats are specified in Appendix D, Section D.6.1.1 and modifications for TOUGHREACT V2.2 are described in Appendix F.

IV.4 Outputs: Data, Records, Files, Screen Displays, and Printed Reports

The software shall produce a print output file electronically that may be printed or plotted. Output files are described in Appendix D, Section D.6.1.2. Modifications for TOUGHREACT V2.2 are described in Appendix F.

IV.5 Output Names, Functional Descriptions, and Sources

The output file name, functional description and source of the output files are described in Appendix D, Section D.6.1.2. Modifications for TOUGHREACT V2.2 are described in Appendix F.

IV.6 Output Frequency and Format

The frequency of output shall vary according to the input options and type of output file. The formats are specified in Appendix D, Section D.6.2. Modifications for TOUGHREACT V2.2 are described in Appendix F.

V. USER INTERFACES

V.1 Error Reporting, Interpretation Standards, and Anticipated Error Conditions

A description of error reporting, interpretation standards, and anticipated error conditions is in Appendix E, Section E.7, for TOUGHREACT V2.1. There will be no modifications for TOUGHREACT V2.2.

V.2 Syntax of Invocation Commands for Script, Batch, and Command Files

There are to be no script, batch, or command files. The software is to be run under Unix by typing the name of the executable file.

V.3 Dialogs

There are to be no dialogs in this software.

V.4 Online Help

There is to be no online help in this software.

V.5 System Messages to Users

The code shall notify the user of errors through error messages on the output file run_log.dat. See Appendix D, Section D.6.1.2, for a description of output file run_log.dat, and Appendix E, Section E.7, for a description of error messages.

VI. SYSTEM INTERFACES

VI.1 Method of Interfacing to Other Systems

No special system interfaces are to be employed.

VI.2 Volume and Frequency of Data Transfer and Data Transfer Formats

Data shall be transferred only by the code reading from and writing to its own files. The formats are specified in Appendix D, Section D.6.2. Modifications for TOUGHREACT V2.2 are described in Appendix F.

VI.3 Verification of Data Transmitted and Received and Checking Data Content and Structure

Verification of data transmitted from the running program to the disk shall not be necessary. File sizes may vary according to the length of the run, the grid size, etc. Other data transfers are governed by YMP-LBNL-QIP-SV.0, *Control of Electronic Management of Data*.

VII. SECURITY

No special design requirements for security are deemed necessary.

VIII. DATA AND LOGICAL MODEL

VIII.1 Data Accuracy, Range, and Default Setting Constraints

Data accuracy and range shall be affected by the user in the form of data inputs. The data accuracy and range are described in Appendix D, Section D.6.2. There are default settings discussed in this report, but they are not constraints on the software system. Any additional data accuracy, range and default settings for TOUGHREACT V2.2 are described in Appendix F.

VIII.2 Data Flow, Data Stores, and Data Entities

For the data flow see flow chart in Appendix C. For data stores and data entities, see description of inputs and outputs described in Appendix D, Section D.6.2, and Section III above. Externally imposed time delays are not allowed.

References:

- Pruess, K., A. Simmons, Y. S. Wu, and G. Moridis, *TOUGH2 Software Qualification*, report LBL-38383, Lawrence Berkeley National Laboratory, Berkeley, California, 1996. MOL.19960610.0010-0020.
- Spycher, N., Sonnenthal, E., Ahlers, R., and Xu, T., *TOUGHREACT V2.1 Software Qualification*, 1999. MOL.20000216.0113
- Wu, Y.S., Haukwa, C., and Mukhopadhyay S., *TOUGH2 V1.4 and T2R3D V1.4: Verification and Validation Report and User's Manual*, Rev 00, 1999. MOL 20000216.0111
- Xu, T., and Pruess, K., *Coupled Modeling of Non-isothermal Multi-phase Flow, Solute Transport and Reactive Chemistry in Porous and Fractured Media: 1. Model Development and Validation*, Lawrence Berkeley Laboratory Report LBNL-42050, 1998. TIC: 243735.

APPENDIX A CONVERSION PLAN

This section does not apply to this software because there is no conversion to be done.

Appendix B: Hardware, Operating System, And Compiler Requirements for TOUGHREACT V2.2 Excerpted from from Spycher et al., 1999, Section 2.1

TOUGHREACT V2.1 has been tested on the following platforms:

<u>Platform</u>	<u>Operating System</u>	<u>Compiler</u>
SUN Ultrasparc systems with single or multiple processors	Solaris SUNOS 5.5.1	SUN f77 (optimization: -O3) Incorrect results are likely if a more aggressive optimization is attempted.
DEC Alpha	DEC-UNIX	DEC F77 with following options: - O4 -warn noalignments -fpe4 -check overflow -assume backslash -check no power -check underflow

Appendix C: Software Structure Requirements for TOUGHREACT V2.2 Excerpted from Xu and Pruess (1998), Section 3

C.3 Model implementation

C.3.1 Solution method

The coupled model is implemented by introducing multi-component transport and reactive chemistry into the framework of the existing non-isothermal multi-phase flow computer code TOUGH2, resulting in the general reactive chemical transport code TOUGHREACT. Our model uses a sequential iteration approach. The flow and transport in geologic media is based on space discretization by means of integral finite differences. An implicit time-weighting scheme is used for flow, transport, and geochemical reaction. An improved equilibrium-kinetics speciation model for simulating water-rock-gas interaction is employed. In this reaction model, equilibrium and kinetics are solved simultaneously by Newton-Raphson iteration, which allows a larger time step to be applied. In addition, the Newton-Raphson iteration scheme has been modified such that the unknowns are the relative concentration increments as opposed to their absolute values. By doing this, the modified Jacobian matrix is symmetric and better conditioned, improving convergence.

The flow chart of the computer simulator TOUGHREACT is presented in Figure 1. The non-isothermal multiphase flow equations are solved first, and the resulting fluid velocities and phase saturations are used for transport simulation. Transport in the liquid phase is treated in terms of total dissolved concentrations. In addition, if gaseous species are present, the transport is solved in terms of their partial pressures. The resulting concentrations and partial pressures from the transport calculation are substituted into the chemical reaction model. The temperature distribution obtained from the solution of the multiphase flow equations is used to update physical and chemical parameters. The chemical transport equations are solved component by component, whereas the reaction equations are solved on a grid block basis. The transport and reaction equations are solved iteratively until convergence.

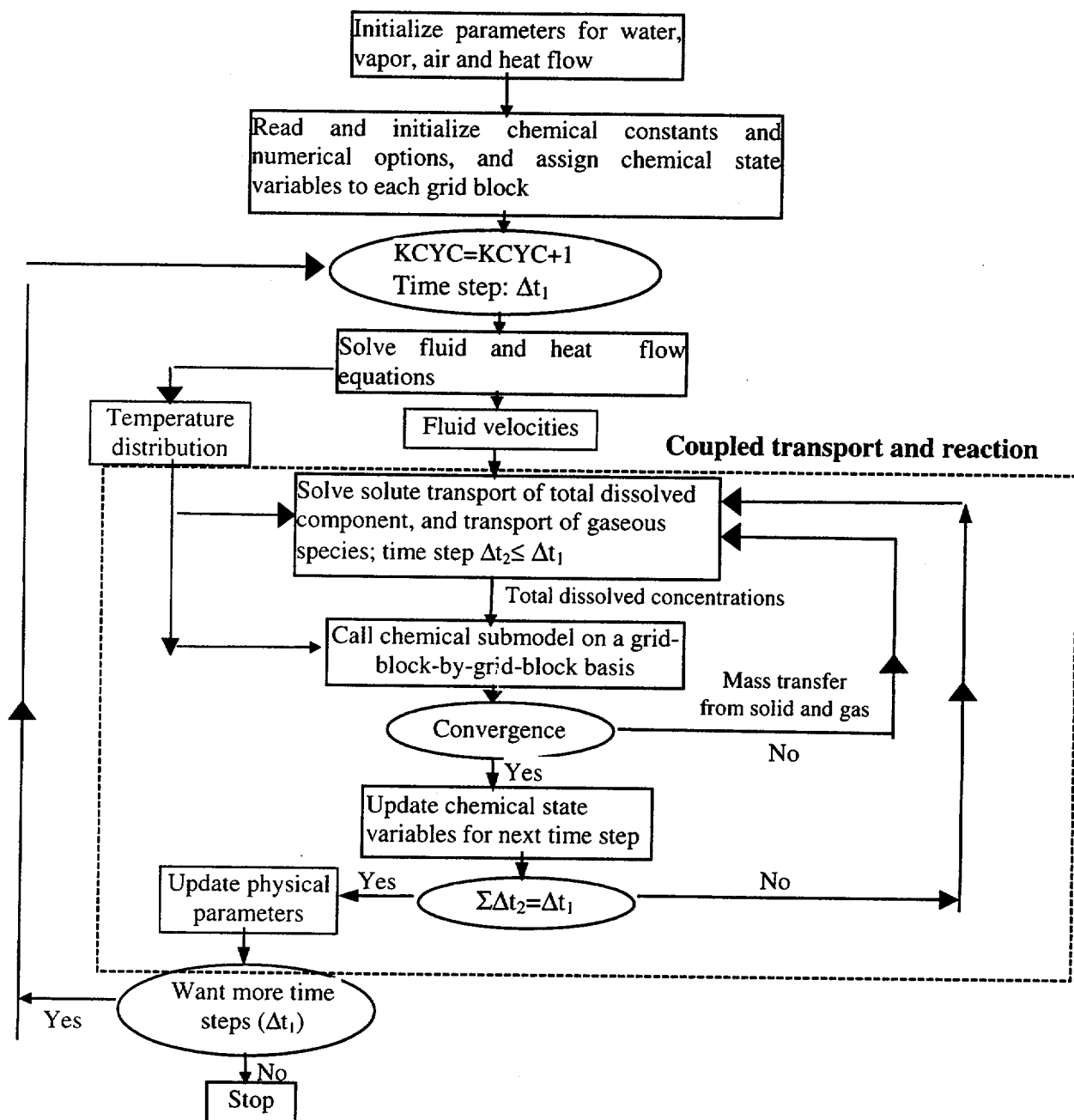


Figure 1. Flow chart of coupled model TOUGHREACT for non-isothermal multiphase flow, solute and reactive chemistry.

C.3.2 Quasi-stationary states

Reactive chemical transport may occur on a broad range of time scales. After a brief transient evolution, reactive systems may settle into a "quasi-stationary state" [QSS], during which aqueous concentrations of all chemical species remain essentially constant. Dissolution of primary and precipitation of secondary minerals proceed at constant rates. In fact, no complex calculations are necessary and only abundances of mineral phases need to be updated. This state terminates when one or more minerals dissolve completely at any of the grid blocks. A relative concentration change, δ_C , and a relative dissolution (or precipitation) rate change, δ_r , are used to monitor attainment of the QSS conditions,

$$\delta_C = \max_{\substack{\text{all components} \\ \text{all grid blocks}}} \left| \frac{C^{k+1} - C^k}{C^k} \right| < \epsilon_C \quad (1a)$$

$$\delta_r = \max_{\substack{\text{all minerals} \\ \text{all grid blocks}}} \left| \frac{r^{k+1} - r^k}{r^k} \right| < \epsilon_r \quad (1b)$$

where k is the transport time step index, C are dissolved component concentrations, r are dissolution or precipitation rates, and ϵ_C and ϵ_r are the QSS tolerances. This is of considerable practical importance, because substantially larger time steps should be possible during periods where a QSS is present.

C.3.3 Time stepping

The solution of the nonlinear reactive transport equations involves several iterative procedures. Each iterative procedure is performed until convergence is achieved. Convergence is very sensitive to the initial estimates of concentrations. At a given time step, initial concentrations are taken equal to those computed at the previous time step. For the first time step, the estimates are taken from the initial concentrations which in some cases may be very different from the true solution. In some cases such as redox reactions a very small time increment may be required in order to give a close initial estimate. Time steps can be increased gradually up to a maximum value when approaching a QSS. An automatic time stepping scheme is implemented in TOUGHREACT. Three time step levels are used. The global time step, Δt_1 , is controlled by the solution of the flow equations. During a time interval of Δt_1 , depending on convergence, multiple steps Δt_2 , with $\sum \Delta t_2 = \Delta t_1$ can be used for solution of transport. Similarly, $\sum \Delta t_3 = \Delta t_2$ (where Δt_3 is the reaction time step) can be used for reaction calculations. The Δt_3 pattern may be different from grid block to grid block depending on the convergence behavior of the local chemical reaction system. For example, at the redox front a small Δt_3 may be required.

C.3.4 Model capabilities

The main features of the TOUGHREACT computer model are listed in Table 5. General multiphase flow and chemical transport conditions are considered. The model can take into account any number of species present in liquid, gas and solid phases. It is well suited for reactive chemical transport in variably saturated porous and fractured media under non-isothermal multiphase flow conditions.

Table 5. Main features of the TOUGHREACT simulator for the coupled modeling of subsurface non-isothermal multiphase flow, solute transport and chemical reactions.

Term	Feature
Dimensionality	1-D, 2-D, or 3-D
Space discretization	Integral finite difference
Time integration	Implicit
Medium	Porous and fractured media; double porosity and dual

	permeability
Heterogeneity	Physical and chemical
Processes for non-isothermal multiphase flow	Fluid flow in both liquid and gas phases occurs under pressure, viscous and gravity forces; capillary pressure, vapor adsorption and vapor pressure lowering effects for the liquid phase; heat flow by conduction, convection and diffusion
Processes for transport of aqueous and gaseous species	advection and diffusion
Linear equation solver for flow and transport	MA28 (LU direct) or preconditioned conjugate gradient (iterative)
Equilibrium reactions	Aqueous complexation, acid-base, redox, mineral dissolution/precipitation, gas dissolution/exsolution, cation exchange, and surface complexation
Kinetics	Mineral dissolution/precipitation
Chemical data base	Modified EQ3/6 data base

Appendix D: General Overview of Input and Output for TOUGHREACT V2.2 from Spycher et al., 1999, Section 6

D.6 TOUGHREACT V2.1 Data Input and Output

This section discusses the program data input and output, including file formats, default parameters, and allowable and tolerable ranges of input parameters.

Input/output files and their formats are generally the same for V2.1 as for V1.0, with a few important changes in the formats of files THERMOK.DAT, SOLUTE.INP, and CHEMICAL.INP. Important changes have also been made in the units of mineral volume fractions and surface areas specified in file CHEMICAL.INP. In addition, in V2.1, the name of the THERMOK.DAT file is no longer fixed and needs to be specified in the SOLUTE.INP file. For these reasons, input files of V1.0 are not upward compatible with V2.1, and vice versa.

D.6.1 General Description of Input and Output Files

D.6.1.1 Input Files

Several data files are used to input data into TOUGHREACT V2.1. All but one input file have fixed file names (i.e. names that cannot be specified by the user).

Fixed-name input files

FLOW.INP - *Hydrological and thermal parameters.* These data include rock properties, initial boundary conditions, time-stepping information, initial conditions of gas and liquid saturations, temperature, and pressure, and all other grid data for hydrological and thermal calculations. The format of this file is identical to the regular TOUGH2 v1.11 input file (Pruess et al. 1996).

SOLUTE.INP - *Transport and other run parameters.* This file contains various flags and input parameters for calculations of reactive transport, such as diffusion coefficients, tolerance limits for convergence of transport and chemical iterations, printout flags for mineral and aqueous species, and the configuration of model zones with different chemical compositions (the compositions of each zone, however, are defined in file CHEMICAL.INP).

CHEMICAL.INP - *Chemical parameters and initial water/rock/gas compositions.* This file is used to define the chemical system (i.e. the type and number of aqueous component species, minerals, and gases considered in the simulation). It also includes the initial compositions of water, mineral, and gas zones configured in file SOLUTE.INP, and kinetic data for minerals (rate constants, surface areas, etc.).

User-specified input files

The names of these files must be specified in the SOLUTE.INP input file, and cannot be left blank. The name of the thermodynamic data base file needs to be specified as follows:

THERMOK.DAT (or any other name) - Thermodynamic database for aqueous species, minerals, and gases. This file contains reaction stoichiometries, dissociation constants ($\log(K)$), and regression coefficients of $\log(K)$'s as a function of temperature.

D.6.1.2 Output Files

Several output files are generated by TOUGHREACT V2.1. These include files with fixed file names, and other files with user-specified names.

Fixed-name output files

FLOW.OUT - Output of thermal and hydrological data (no chemical information). This file is identical to a TOUGH2 V1.11 output file, including data on temperature, pressure, liquid and gas saturations, liquid and gas (steam+ air) fluxes, and velocities for all grid blocks of the model.

CHEMICAL.OUT - Echo of chemical input data. This file lists the data read from input files CHEMICAL.INP and THERMOK.DAT, including initial water rock, and gas compositions, equilibrium constants and stoichiometries of chemical reactions, and kinetic data.

SOLUTE.OUT - Echo of transport and other run parameters. This file lists the data read from input file SOLUTE.INP, including all transport parameters, chemical zone configuration, and other run-specific parameters.

run_log.dat - Log of the simulation progress. This file is constantly updated during a simulation. It lists some run input parameters and all run-related messages, including error messages (Section 7). It also contains mass balance and other data whose printout is enabled by setting specific MOP parameters to non-zero values in the FLOW.INP file

chdump.dat - chemical speciation data. This file contains results of geochemical speciation calculations for each initial water composition input into the model. It also lists these data for grid nodes where chemical convergence fails. The results are computed for an original 1 liter of liquid. They include a printout of chemical mass balances (total mass balance and aqueous species mass balance) which must always remain zero or very small. For debugging purposes or for small grids, if the flag ICHDUMP in the SOLUTE.INP input file is set equal to 1, geochemical speciation results will be output in the *chdump.dat* file for every grid node at every time step (!). As a precaution to avoid filling up disk space, results of speciation calculations are output only for the first thousand grid nodes and/or time steps, after which the program will abort.

User-specified output files

The names of these files must be specified in the SOLUTE.INP input file, and cannot be left blank. The file names currently specified and the file contents are as follows:

ITER.DAT - Iteration Data. This file lists the number of flow, transport and chemical iterations used to reach convergence at each simulation time step. The number of transport iterations in this file indicate the number of iterations between flow, transport and chemical computations at each time step (not the actual number of iterations required to reach convergence during calculations of aqueous species and gas transport). It is always equal to 1 with this version of the program (no iteration between flow, transport, and chemistry).

TEC_CONC.DAT - Aqueous species plot data. This file contains time (s), grid block coordinates (m), liquid saturation, temperature ($^{\circ}\text{C}$), pH, and aqueous species concentrations ($\text{mol/liter}_{\text{liquid}}$) at all grid blocks for time steps specified in the FLOW.INP file. The number and type of species to output in the TEC_CONC.DAT file is specified by flags in the input file SOLUTE.INP. This file is in a TECPLOT-compatible format, although some editing of this file will be required to import data for dual permeability grids into TECPLOT. Note: there is a field for oxygen fugacity in this file. This field is padded with zero values unless a redox system is considered. However, redox simulations have not been tested with this version of the program.

TEC_MIN.DAT- Mineral and gas plot data. This file contains time (s), grid point coordinates (m), temperature ($^{\circ}\text{C}$), changes in mineral amounts ($\text{mol/m}^3_{\text{medium}}$), and gas pressures (bars) (e.g. CO_2) at all grid blocks for time steps specified in the FLOW.INP file. Currently, mineral amounts are output in moles per cubic meter of rock medium (i.e. matrix or fracture), and gas pressures in bars. The mineral amounts represent the change between the amount at the current time step and the initial amount (at time zero). Negative amounts indicate overall dissolution, and positive amounts overall precipitation. The number of minerals in the TEC_CONC.DAT file are specified by flags in the input file SOLUTE.INP. Gas pressures for all gases specified in the CHEMICAL.INP file are output following the mineral data. This file is in a TECPLOT-compatible format, although some editing of the file will be required to import data for dual permeability grids into TECPLOT.

TIME.DAT - Plot data at specified grid blocks. This file contains grid block identifier, time (days), liquid saturation, temperature ($^{\circ}\text{C}$), pH, aqueous species concentrations ($\text{mol/l}_{\text{liquid}}$), mineral amounts ($\text{mol/m}^3_{\text{medium}}$), and gas pressures (bars) for specific grid blocks and time step intervals, as specified in the input file SOLUTE.INP. This file can be used to generate time-profiles at specified locations. Note: there is a field for oxygen fugacity in this file. This field is padded with zero values unless a redox system is considered. However, redox simulations have not been tested with this version of the program.

D.6.2 Input/Output File Formats and Contents

Required formats for each input file are described below. FORTRAN format notations are used to describe fixed formats (A for strings, I for integers, and F for reals, X for spaces, etc.). Free formats are described using a modified FORTRAN notation, with format letters specified without a length. Free formats for arrays may be preceded by a variable (e.g. `ntmp*F`) which indicates the maximum dimension of the array. These dimensions are generally defined in the `PARAMETER.INC` source file. Character strings in free format must be enclosed in single-quotes. As a reminder, free format string variables below are listed with names between quotes.

D.6.2.1 FLOW.INP - Hydrological and thermal parameters

Format: fixed

No changes since V1.0. The format of this file is identical to the regular TOUGH2 input file (Pruess 1991). However, a few MOP parameters have been assigned new functions, as follows:

- MOP(20) = 0 for reactive transport
 = 1 no reactive transport, but input files with chemical data are read
 > 1 no reactive transport, no chemical data files are read
- MOP(22) > 0 writes the transport coefficient matrix, Darcy velocities, porosities, and other transport data in the *run_log.dat* file during calculations of aqueous species and gas transport. For debugging only.
- MOP(23) > 0 writes source terms, right-hand side, old and new aqueous concentrations, and various other parameters in the *run_log.dat* file during transport calculations. Also outputs the permeability, porosity, and capillary pressure correction factor at each grid block in the *run_log.dat* file. For debugging purposes only.
- MOP(24) =1 does not force at least one transport step to be calculated (can be useful to compute chemical reaction only in single-grid node problems).

D.6.2.2 THERMOK.DAT - Thermodynamic data for aqueous species, minerals, and gases.

Format: free

The format of this file has changed since version V1.0. Mineral molecular weights have been added to the first record of entries for these components. Otherwise, entries in this file have remained unchanged since V1.0. The new file version is not backward compatible with TOUGREACT V1.0, nor is the old file version upward compatible with TOUGHREACT V2.1.

Initial records:

- First record (A,I,ntemp*F): 'DUMMY', NTEMP, (TEMPC(i), i=1,NTEMP)

DUMMY is a label used to describe the data for this record. NTEMP is the number of TEMPC values to read. TEMPC are temperatures (°C) at which the log(K) data are listed in this file. TEMPC values must be listed in order of increasing temperature. Only the first (minimum) and last (maximum) temperatures are used by the program. These values are used to constrain log(K) extrapolation within this temperature range. Log(K)'s are not extrapolated outside this temperature range! For example, if the maximum TEMPC is 150 °C but the computed system temperature is 250 °C, log(K)'s will be extrapolated only to 150 °C (i.e. the geochemical speciation will be computed at 150 °C, not 250 °C)! Therefore, users must make sure that simulation temperatures are within the range of thermodynamic data temperatures.

Component species records. These data immediately follow the initial file record, as follows:

- Each record (A,2F): 'NAME', A0, Z

For each component species, NAME is the name, A0 the Debye-Huckel a0 parameter (ion effective or hydrated radius), and Z the ion electric charge. The maximum length of NAME is 20. The maximum number of records is *mpri* (defined in source file PARAMETE.INC)

- Last record must be (A,2F): 'null' 0.0 0.0 (where 'null' is the actual string in quotes)

Derived species records. These must directly follow the component species records. The data for each derived species is given by 3 records, which are as follows:

- 1st record (A,2F,I,mpri*(F,A)): 'NAME', A0, Z, NCPS, (STQS(i), 'NAM(i)', i=1,NCPS)

NAME, A0, and Z are as defined for the component species. NCPS is the number of component species defining the derived species. STQS contains the stoichiometric coefficient of component species NAM included in the dissociation reaction of the derived species (negative and positive values for reactants and products, respectively). The maximum value for NCPS is *mpri* (define in source file PARAMETE.INC).

- 2nd record (A,ntemp*F): 'NAME', (DUMMY(i), i=1,ntemp)

NAME is as defined above. DUMMY contains the dissociation constants (log(K) in base 10) for the given reaction at each discrete temperature listed in the initial file record. These data

are skipped on input, because all $\log(K)$ values are computed as a function of temperature using the regression coefficients that follow (below). The discrete $\log(K)$ values should, however, always be included in the file to provide for an easy reference of the data.

- 3rd record (A,5E): 'NAME', (AKCOES(*maq*_x,*i*), *i*=1,5)

NAME is as defined above. AKCOES contains regression coefficients *a*, *b*, *c*, *d*, and *e* to calculate $\log(K)$ as a function of temperature (within the range of temperatures listed on the first record of the file) such that $\log(K) = a \cdot \ln(T_k) + b + c \cdot T_k + d/T_k + e/T_k^2$, where *T_k* is absolute temperature (K), and *log* and *ln* stand for base-10 and natural logarithms, respectively.

The maximum total number of derived species allowed in the program is *maq*_x (defined in source file PARAMETE.INC). After the data for all derived species have been listed (three records per species), the last record to end the derived species data must be as follows:

- Last record (A,2F,I): 'null' 0. 0. 0 (where 'null' is the actual string in quotes)

The mineral records. These must directly follow the derived species records. The data for each mineral is given by 3 records, which are as follows:

- 1st record (A,2F,I,*mpri**(2F,A)): 'NAME', MOLWT, VMIN, NCPM, (STQM(*i*), NAM(*i*)), *i*=1,NCPM)

NAME, MOLWT and VMIN are the name of the mineral, its molecular weight (g/mol) and molar volume (cm³/mole), respectively. NCPM is the number of component species defining the mineral. STQM contains the stoichiometric coefficient of component species NAM in the dissociation (hydrolysis) reaction of the mineral (negative and positive values for reactants and products, respectively). The maximum value for NCPM is *mpri* (defined in source file PARAMETE.INC).

- 2nd record (A,*ntmp**F): 'NAME', (DUMMY(*i*), *i*=1,*ntmp*)

NAME is as defined above. DUMMY is the dissociation constants ($\log(K)$ in base 10) for the given reaction at each discrete temperature listed in the first record of the file. These data are skipped on input, because all $\log(K)$ values are computed as a function of temperature using the regression coefficients that follow (below). The discrete $\log(K)$ values should, however, always be included in the file to provide for an easy reference of the data.

- 3rd record (A,5E): 'NAME', (AKCOEM(*mmin*,*i*), *i*=1,5)

NAME is as defined above. AKCOEM contains regression coefficients a , b , c , d , and e to calculate $\log(K)$ as a function of temperature (within the range of temperatures listed on the first record of the file) such that $\log(K) = a \cdot \ln(T_k) + b + c \cdot T_k + d/T_k + e/T_k^2$, where T_k is absolute temperature (K), and \log and \ln stand for base-10 and natural logarithms, respectively.

The maximum total number of minerals allowed in the program is *mmin* (defined in source file PARAMETE.INC). After the data for all minerals have been listed (three records per mineral), the last record to end the mineral data must be as follows:

- Last record (A,2F,I): 'null' 0. 0 (where 'null' is the actual string in quotes)

The gas records. These must directly follow the mineral records, and have a format identical to that of minerals. The data for each gas is given by 3 records as follows:

- 1st record (A,F,I,mpri*(F,A)): 'NAME', DUMMY, NCPG, (STQG(i), 'NAM(i)', i=1,NCPG)

NAME is the name of the gas. DUMMY is molar volume but this value is not used (the molar volume of gases are computed by the program assuming the ideal gas law). NCPG is the number of component species defining the gas. STQG contains the stoichiometric coefficient of component species NAM in the dissociation reaction of the gas (negative and positive values for reactants and products, respectively). The maximum value for NCPG is *mpri* (defined in source file PARAMETE.INC).

- 2nd record (A,ntmp*F): 'NAME', (DUMMY(i), i=1,ntmp)

NAME is as defined above. DUMMY is the dissociation constants ($\log(K)$ in base 10) for the given reaction at each discrete temperature listed in the first record of the file. These data are skipped on input, because all $\log(K)$ values are computed as a function of temperature using the regression coefficients that follow (below). The discrete $\log(K)$ values should, however, always be included in the file to provide for an easy reference of the data.

- 3rd record (A,5E): 'NAME', (AKCOEG(mgas,i), i=1,5)

NAME is as defined above. AKCOEG contains regression coefficients a , b , c , d , and e to calculate $\log(K)$ as a function of temperature (within the range of temperatures listed on the first record of the file) such that $\log(K) = a \cdot \ln(T_k) + b + c \cdot T_k + d/T_k + e/T_k^2$, where T_k is absolute temperature (K), and \log and \ln stand for base-10 and natural logarithms, respectively.

The maximum total number of gases allowed in the program is *mgas* (defined in source file PARAMETE.INC). After the data for all gases have been listed (three records per gas), the last record to end the mineral data must be as follows:

- Last record (A,F,I): 'null' 0. 0 (where 'null' is the actual string in quotes)

The adsorbed species records (surface complexes). This feature of TOUGHREACT has not been tested in V2.1 and therefore no data should be entered, except for the termination record:

- Last record (A,F,I): 'null' 0. 0 (where 'null' is the actual string in quotes)

D.6.2.3 SOLUTE.INP - Transport and other run parameters

Format: fixed

The format of this file has changed since version V1.0. The name of the thermodynamic database must now be entered in this file, and a new factor is read in the fifth record. In other respects, entries in this file have remained unchanged since V1.0. The new file version is not backward compatible with TOUGHREACT V1.0, nor is the old file version upward compatible with TOUGHREACT V2.1.

- 1st record (A76): TITLE

Run title

- 2nd record (): skipped on input; can be used to define the data that follows
- 3rd record (3I5,F5.3,3I5): ISPIA, INIBOUND, ISOLVC, ICOUR, NGAS1, ICHDUMP, KCPL

ISPIA is a flag to define the sequential iterative approach between computations for transport and for chemistry, as follows:

- 0 sequential iteration between transport and chemistry (if MAXITPTR > 1)
- 1 partial sequential iteration (do not use, not tested!)
- 2 no sequential iteration (fully explicit reaction source terms)
- 3 sequential iterations between transport, chemistry, and flow (do not use, not tested!)

INIBOUND is a flag for enabling the use of different recharge water compositions (0=disabled, 1=enabled)

ISOLVC is a flag for the type of transport solver to use. Always use ISOLVC = 3 (default) with this version of the program. Other options have not been tested.

RCOUR is both a variable and a flag to limit the time step size. RCOUR > 0.0 limits the maximum time step size to RCOUR x Courant Number. This option is disabled if RCOUR = 0.0.

ICHDUMP is a flag to enable printing of chemical speciation results at each grid node and each time step (0 = disabled, 1=enabled). It should be enabled for debugging purposes only, or to track small problems. If this option is enabled, the program will abort after outputting speciation results for the first 1000 grid nodes and/or time steps, to avoid accidentally filling up disk space.

KCPL is a flag to enable coupling porosity, permeability, and capillary pressure changes as described in Section 3 (0 = disabled, 1=enabled).

- 4th record (): skipped on input; can be used to define the data that follows.
- 5th record (3E10.4): SL1MIN, D1MIN, STIMAX, CNFACT

The first three parameters are used for skipping the chemical solver (i.e. skip geochemical speciation calculations) at grid nodes where conditions of saturation, inter grid block distance, or ionic strength are outside of the valid ranges of the model. The geochemical calculations are skipped at grid blocks where: the liquid saturation is less than SL1MIN; the minimum distance to the center of any adjacent block is less than D1MIN; or the stoichiometric ionic strength is more than STIMAX. For typical boiling simulations, use SL1MIN less or equal to 10^{-3} . Set D1MIN = 0.0 (disabled) unless absolutely necessary. With this program version, STIMAX should not be greater than 2.0 (preferably 1.0) because of limitations regarding the calculation of activity coefficients at elevated ionic strength.

CNFACT is a weighting factor for mineral and gas reaction source terms in the transport equations (1.0 = fully implicit source terms, 0.0 = fully explicit source terms). This parameter has an effect only if sequential iterations are enabled (ISPIA = 0 or 1). In this program version, CNFACT always defaults to 1.0 if a non-zero value is input (implicit only). Simulations with CNFACT = 0.0 using sequential iterations will produce the same results as simulations without sequential iterations (explicit source terms) but requires increased computing time and therefore should be avoided.

- 6th record (): skipped on input; can be used to define the data that follows

- 7th record (A20,A60): OUTITER, TITLE
Name file for output of iteration data (ITER.DAT), and any title
- 8th record (A20,A60): OUTPLOT, TITLE
Name of file for output of aqueous species plot data (TEC_CONC.DAT) and any title.
- 9th record (A20,A60): OUTSOLID, TITLE
Name of file for output of mineral and gas plot data (TEC_MIN.DAT) and any title
- 10th record (A20,A60): OUTTIME, TITLE
Name of file where to output time profile plot data (TIME.DAT) and any title
- 11th record (): skipped on input; can be used to define the data that follows
- 12th record (2F10.0,2E10.3): WTIME,WUPC,DIFUN,DIFUNG

WTIME is an interface weighting factor. Always set WTIME = 1 (other values not tested). WUPC is an upstream weighting factor. Always set WUPC = 1 for full upstream weighting (other values not tested). DIFUN and DIFUNG are the products of the diffusion coefficient and tortuosity of the medium (m^2/sec) for aqueous species and gases, respectively.

- 13th record (): skipped on input; can be used to define the data that follows
- 14th record (I5,E10.3,I5,E10.3,I5,E10.3,I5,E10.3): MAXITPTR, TOLTR, MAXITPCH, TOLCH, MAXITPAD,TOLAD

MAXITPTR is the maximum number of iterations for solving the sequential transport/chemistry iterative procedure. If MAXITPTR=1, no sequential iteration will take place even if ISPIA \neq 2. TOLTR is the relative concentration tolerance for convergence during sequential iterations (difference between the concentrations at the current and previous and sequential iterations, divided by the arithmetic average of these concentrations).

MAXITPCH is the maximum number of iterations for solving geochemical speciation (typically 200). TOLCH is the relative concentration tolerance for solving geochemical speciation (concentration change from the previous iteration to the current one, divided by the current concentration). Set TOLCH $\leq 10^{-4}$.

MAXITPCH and TOLAD are maximum number of iterations and tolerance for solving adsorption. This version of the program was not tested with surface complexes. These

parameters are not used if no adsorption species are entered in the CHEMICAL.INP file (do not enter any).

- 15th record (): skipped on input; can be used to define the data that follows
- 16th record (7I5): NWTI, NWNOD, NWCOM, NWMIN, IWCOMT

NWTI is a flag for writing data in the TIME.DAT file. Data will be written at every NWTI time steps.

NWNOD is the total number of grid blocks at which to output data in the TIME.DAT file. It should correspond to the number of nodes listed on record 18.

NWCOM is the total number of aqueous species for which to output concentrations in the TIME.DAT and TEC_CONC.DAT files. It should correspond to the number of species listed on record 20.

NWMIN is the total number of minerals for which to output amounts in the TIME.DAT and TEC_MIN.DAT files. It should correspond to the number of species listed on record 22.

IWCOMT is a flag to indicate the type of aqueous species concentration to output in the TIME.DAT and TEC_CONC.DAT files. IWCOMT=0 for concentrations of individual aqueous species. IWCOMT=1 for total aqueous concentrations (i.e. the sum of concentrations of a component species and all derived species that contain this component).

- 17th record (): skipped on input; can be used to define the data that follows
- 18th record (15A5): ELEMW(i), i = 1,NWNOD

ELEMW contains the list of grid node names for which to output data in the TIME.DAT file. If NWNOD is greater than 15, an additional record must be inserted, which offsets all following records. The maximum number of node names is 30. Leave the record blank if NWNOD = 0.

- 19th record (): skipped on input; can be used to define the data that follows
- 20th record (15I5): IWCOM(i), i = 1,NWCOM

IWCOM contains the list of species indexes for which to output data in the TIME.DAT file and TEC_CONC.DAT file. The indexes correspond to the species order in the list of

component species entered in CHEMICAL.INP. For derived species (if IWCOMT = 0) indexes above those of component species can be determined from the chdump.dat file by aborting a run after the first time step. If NWCOM is greater than 15, an additional record must be inserted, which offsets all following records. The maximum allowed number of species indexes is 30. Leave this record blank if NWCOM=0.

- 21th record (): skipped on input; can be used to define the data that follows
- 22th record (20I5): IWMIN(i), I = 1,NWMIN

IWCOM contains the list of mineral indexes for which to output data in the TIME.DAT file and TEC_CONC.DAT file. The mineral indexes correspond to the order in the list of minerals entered in CHEMICAL.INP. The maximum allowed number of mineral indexes is 20. Leave this record blank if NWMIN=0.

- 23th record (): skipped on input; can be used to define the data that follows
- 24th record (6I5): IZIWDF, IZBWDF, IZMIDF, IZGSDF, IZADDF, IZEXDF

The first four parameters are the default initial waters, boundary waters, mineral zones, and gas zones, respectively, to assign to each grid node. Values of IZIWDF and IZBWDF must each correspond to one of the water composition indices (IWTYPE) specified in CHEMICAL.INP. Similarly, values of IZMIDF and IZGSDF must correspond to one of the mineral and gas zone indices (IMTYPE and IGTTYPE, respectively) specified in CHEMICAL.INP. The last two parameters are not used with this version of the program and must be set to zero.

- 25th record (): skipped on input; can be used to define the data that follows
- Next records (one per grid block) (A5,8I5): EL, NSEQ, NADD, IZIW, IZBW, IZMI, IZGS, IZAD, IZEX

These records are used to assign initial waters, boundary waters, mineral zones, and gas zones to specific grid nodes of the model (maximum allowed is *mnod*, defined in source file PARAMETE.INC). These records can be omitted (in this case all grid blocks are assigned the defaults specified on record 24). Parameters as follows for each grid block:

EL	grid node name
NSEQ	= 0 (always, with this version of the program)
NADD	= 0 (always, with this version of the program)

IZIW	initial water index (one of the IWTYPE values in CHEMICAL.INP)
IZBW	boundary water index (one of the IWTYPE values in CHEMICAL.INP)
IZMI	mineral zone index (one of the IMTYPE values in CHEMICAL.INP)
IZGS	gas zone index (one of the IMTYPE values in CHEMICAL.INP)
IZAD	= 0 (always, with this version of the program)
IZEX	= 0 (always, with this version of the program)

- End record for zone assignments (A5,8I5): (must be blank)
- Next record (): skipped on input; can be used to define the data that follows
- Next record (A5,5E10.3): EL, DISG(i), (PFUGB(i,ig), ig = 1,NGAS)

This record is to specify nodes connected to a gas supply (i.e. atmosphere). This option has not been tested and this record must be left blank.

Final file record (optional): end ('end' is the actual string, without quotes)

D.6.2.4 CHEMICAL.INP - Chemical parameters and initial water/rock/gas compositions

Format - free

The units of mineral surface areas and initial volume fractions specified in this file have changed since version V1.0. Also, a variable was added to the record specifying parameters for minerals precipitation rates. Therefore, this file is not backward compatible with V1.0 (or vice versa).

Initial records

- 1st record (A): 'LABEL'
- 2nd record (A): 'LABEL'
- 3rd record (A): 'LABEL'

LABEL is a character string (maximum length 100) currently used to define the run title (1st and 2nd record) and to describe the data that follows (3rd record).

Component species records. These define the chemical system in the simulation, and must be entered directly following the initial records. One record must be entered per component species, in any order, as follows:

- Initial record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

- Each following record (A): 'NAPRI'

NAPRI is the name of each selected component species (20 characters maximum). These names must match exactly the names of component species in the THERMOK.DAT file.

The maximum total number of component species allowed in the program is *mpri* (defined in source file PARAMETE.INC). After the selected component species have been listed (one record per species), the last record to end the component species list must be as follows:

- Last record (A): '*' (where '*' is the actual string in quotes)

The program selects automatically in the file THERMOK.DAT all derived species that contain the selected component species.

Mineral records. These contain the minerals desired in the simulation, and must directly follow the component species records. Minerals can be entered in any order as long as the minerals at equilibrium precede those under kinetic constraints. The specified minerals consist of reactants and any possible products. Their names must match exactly the names of minerals in the THERMOK.DAT file. Also, all component species used in the mineral reaction stoichiometry in the THERMOK.DAT must be included in the list of component species above.

Minerals can be specified to react either at equilibrium with the aqueous phase, or under kinetic constraints. Minerals with identical stoichiometries (i.e. quartz and cristobalite) cannot both be specified at equilibrium (use kinetic constraints in this case). Minerals specified to be at equilibrium are equilibrated with the solution only if these are calculated to be supersaturated (no forced equilibrium). If undersaturated, these minerals are assumed to be totally dissolved and remain as possible products (inactive bystanders) until they reach saturation.

- Initial record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

Minerals at equilibrium are defined with one record (per mineral). Minerals under kinetic constraints require one or two more records (per mineral). Minerals at equilibrium must always be listed preceding those under kinetic constraints. The following record(s) must be specified for each mineral:

- First mineral record (A, 2I): 'NAMIN', IKIN, IDISPRE

NAMIN is the mineral name (20 characters maximum). IKIN is a flag for the type of mineral: 0 for minerals at equilibrium, and 1 for those under kinetic constraints. IDISPRE is a flag for the type of kinetic constraints: 1 for dissolution only, 2 for precipitation only, and 3 for both (mineral can either precipitate or dissolve). Always set IDISPRE = 0 if IKIN = 0 and IDISPRE > 0 if IKIN = 1.

- 2nd mineral record, only if IKIN > 0 (F,I,6F): RKF, IDEP, CK1, CK2, EA, ACFDISS, BCFDISS, CCFDISS (mineral dissolution rate parameters)

RKF is the coefficient a in the expression $k = a \cdot \exp(EA/RT)$, where k is the rate constant (in mol/m²/sec), EA is the activation energy in kJ/mol, R the ideal gas constant (8.3144·10⁻³ kJ/°K/mol), and T absolute temperature (°K). The form of the rate law is:

$$\text{rate} = k \cdot S \cdot (1 - (Q/K)^m)^n$$

where the rate is in mol/sec/kg_{H2O}, Q/K is the mineral saturation index, and S the effective surface area (in m²/kg_{H2O}) specified later.

IDEP is a flag for rate dependence on pH. Always use IDEP=0 with this version of the program (no pH dependence).

CK1 and CK2 are the exponents n and m , respectively in the above rate expression. Always use CK1=1 and CK2=1 with this version of the program.

ACFDISS, BCFDISS, and CCFDISS should be set to zero, unless a different form of rate constant dependence with temperature is desired. This alternate form is: $\log(k) = a + b \cdot T + c/T$, where T is absolute temperature in K and \log is in base 10. To enable this option, RKF must be set to 1.0, EA must be set to zero and ACFDISS, BCFDISS, and CCFDISS must be specified as the coefficients a , b , and c , respectively, in the above expression.

- 3rd mineral record, only if IKIN > 0 and IDISPRE = 2 or 3 (F,I,6F): RKPREC, IDEPREC, CK1PREC, CK2PREC, EAPREC, ACFPREC, BCFPREC, CCFPREC, RNUCL

(mineral precipitation rate parameters)

These input parameters, except for the last one, are listed in the same order and have the same functions as those described above for mineral dissolution (2nd record), except that the parameters apply to mineral precipitation instead of dissolution.

RNUCL contains the initial volume fraction ($V_{\text{mineral}}/V_{\text{solid}}$) to be assumed for calculating initial effective surface area (Equation 3-4) if the mineral is not present at the start of a simulation but precipitates as a new reaction product. If zero, RNUCL is assumed to be 10^{-5} .

The maximum total number of minerals allowed in a simulation is *mpri* (defined in source file PARAMETE.INC). After the selected minerals have been listed (one record per mineral), the last record to end the minerals list must be as follows:

- Last record (A,2I): '*' 0 0 (where '*' is the actual string in quotes)

Gas records. These contain gas species desired in the simulation (besides H₂O, which is treated independently in the flow routines). The gas names must match exactly the names of gases in the THERMOK.DAT file. Also, all component species used in the gas reaction stoichiometry in the THERMOK.DAT must be included in the list of component species above. The gas records must be entered directly following the mineral records, in any order, as follows:

- Initial record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

- Each gas record (A): 'NAGAS'

NAGAS is the gas name (20 characters maximum). The maximum total number of gases allowed in the program is *mgas* (defined in source file PARAMETE.INC). The last record to end the gas list must be as follows:

- Last record (A): '*' (where '*' is the actual string in quotes)

Adsorbed species records (surface complexes). These directly follow the gas records. This feature of TOUGHREACT has not been tested with surface complexes in V2.1 and therefore no data should be entered, except for a termination record:

- Last record (A): '*' (where '*' is the actual string in quotes)

Initial, boundary, and recharge water compositions. These data directly follow the gas records, and are headed by two label records as follows:

- Initial record (A): 'LABEL'
- Next record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

- Next record (3I): NIWTYPE, NBWTYPE, NRWTYPE

These parameters indicate the number of different sets of initial water compositions, boundary water compositions, and recharge water compositions to read in, respectively. The maximum number of different water compositions is *mnel* initial waters, *mbound* boundary waters, and *mrech* recharge waters (these parameters are defined in source file PARAMETE.INC).

Sets of water compositions follow. A total of NIWTYPE + NBWTYPE + NRWTYPE sets must be entered. These sets must be listed in order of first, initial waters, then boundary waters, then recharge waters. Each water composition set has the following format:

- 1st record (I,F): IWTYPE, TC2

IWTYPE is an index for the water composition set. It must be 1 for the first set, 2 for the second set, and so on, but must be reset to 1 at the start of each different water type (i.e. for the first initial, first boundary, and first recharge water). This index is used in file SOLUTE.INP to assign water compositions to grid blocks. TC2 is the initial temperature of the water (°C) at which the initial geochemical speciation is calculated. This temperature does not affect flow/thermal calculations and will be overwritten by the initial system temperature specified in the FLOW.INP file. If TC2 is much different than the latter temperature, the program may have difficulty in reaching convergence when computing geochemical speciation during the first time step.

- 2nd record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

- Following records (one record per species) (A,I,2F,A,I): 'NAME', ICON, CGUESS, CTOT, 'DUM', ICTOT2

NAME is the name of component species as specified in the component species list in the first part of the file, but not necessarily in the same order. ICON is a flag which indicates the type of concentration to be entered. It must be either 1 or 3, as described below.

CGUESS is the trial molality of the component species, for use as a starting point for Newton-Raphson iterations necessary to solve geochemical speciation. CGUESS can take

any value, but should be as close as possible to the actual molality of the species (the calculated solution) or convergence may not occur.

CTOT stores the species concentration (i.e. the analyzed water composition) and takes different input depending on the value of ICON:

If ICON=1: CTOT for all components except water is total concentration (component + derived species) in moles per specified liter of liquid. For water, the amount entered is the volume of liquid in liters (generally one liter so that CTOT for other components is in moles/liter).

If ICON=3: CTOT is the activity of the individual component species (for example 10^{pH} for H^+). Do not use this option for water.

DUM is an unused dummy string variable. Always leave as one or more blanks between quotes.

ICTOT2 is currently unused. Always set $\text{ICTOT2} = 0$

Each water composition set must be terminated with the following record:

- End record (A,I,2F,A,I): `*' 0 0.0 0.0 '' 0`

(where `*` and `''` are a star and a blank string in quotes)

Initial mineral zones. These records directly follow the water composition sets and are headed by two label records as follows:

- Initial record (A): `'LABEL'`
- Next record (A): `'LABEL'`

LABEL is a character string (maximum length 100) used to describe the data that follows.

- Next record (I): `NMTYPE`

This parameter indicates the number of mineral zones to read in (maximum is *mnel*, defined in source file `PARAMETE.INC`).

Records for mineral zones follow. A total of NMTYPE mineral zones must be listed. Each mineral zone has the following format:

- 1st record (I): IMTYPE

This is the index of the mineral zone. It must be 1 for the first zone, 2 for the second zone, and so on. This index is used in file SOLUTE.INP to assign the mineral zones to grid blocks.

- 2nd record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

Each mineral at equilibrium is then defined with one record. Minerals under kinetic constraints require one more record. Minerals can be listed in any order as long as those at equilibrium precede those under kinetic constraints. Mineral spelling and designation (equilibrium or kinetic) must be the same as defined previously. The following record(s) must be specified for each mineral:

- 1st mineral record (each mineral) (A,F,I): 'NAME', VOL2, IKIN4

NAME is the name of the mineral (20 characters maximum). VOL2 is the initial volume fraction of the mineral, excluding liquid (mineral volume divided by total volume of solids). The sum of VOL2's need not add up to 1. The remaining solid volume fraction is considered unreactive. IKIN4 is a flag to indicate whether the mineral is to be treated at equilibrium (IKIN4=0) or under kinetic constraints (IKIN4=1). IKIN4 must be consistent with previously defined IKIN for each mineral.

- 2nd mineral record, only if IKIN4 = 1 (2F): RAD, AMIN

RAD is the grain radius, which is currently not used (set to zero or any value). AMIN is the mineral surface area in [cm²/g] (i.e. square centimeters per gram of mineral). AMIN is not the effective reactive surface area (see Section 3).

After all the minerals in the zone have been listed (one record per mineral), the last record to end the mineral zone must be as follows:

- Last record (A,F,I): '*' 0.0 0 (where '*' is the actual string in quotes)

Initial gas zones. These records directly follow the mineral zones and are headed by two label records as follows:

- Initial record (A): 'LABEL'
- Next record (A): 'LABEL'

'LABEL' is a character string (maximum length 100) used to describe the data that follows.

- Next record (I): NGTYPE

This parameter indicates the number of gas zones to read in (maximum is *mmel*, defined in source file PARAMETE.INC).

Records for each gas zone follow if NGTYPE > 0. A total of NGTYPE gas zones must be entered. Each gas zone has the following format:

- 1st record (I): IGTTYPE

This is the index of the gas zone. It must be 1 for the first zone, 2 for the second zone, and so on. This index is used in file SOLUTE.INP to assign the gas zones to grid blocks.

- 2nd record (A): 'LABEL'

'LABEL' is a character string (maximum length 100) used to describe the data that follows.

- Each following record (one per gas) (A,F): 'NAME', CGNOD

NAME is the name of the gas (20 characters maximum). Gas names must be the same as defined previously (in any order). CGNOD is the gas initial partial pressure (in bars) in the fraction of pore spaces that are not filled with water. If this pressure is not the same as the partial pressure reflecting the initial composition of the pore water, a new initial partial pressure will be calculated to reflect chemical equilibrium between pore gas and pore water. If the initial gas fraction in pore spaces is small, CGNOD has little effect on calculation results. However, if the initial gas fraction is large, the value of CGNOD may contribute significantly to the total initial mass of the gas in the system.

After all the gases in the zone have been listed, the last record to end the gas zone must be as follows (also needed if NGTYPE=0):

- Last record (A,F): '*' 0.0 (where '*' is the actual string in quotes)

Initial adsorption zones. This feature of TOUGHREACT was not tested in V2.1. Therefore, these data must be left blank. However, for the program to finish reading the file correctly, several records must be included directly following the gas zone data:

- Initial record (A): 'LABEL'
- Next record (A): 'LABEL'

LABEL is a character string (maximum length 100) used to describe the data that follows.

- Next record (I): NDTYPE

Set NDTYPE = 0 (no surface complexes).

- Next record (A): 'LABEL' (as above)

Last two records of file. These must be present to avoid input errors:

- Last before final record (A): 'LABEL' (as above)

LABEL is any character string (maximum length 100)

- Final record in file (A): 'end' (where 'end' is the actual string in quotes).

Appendix E: Error Messages and Error Handling for TOUGHREACT V2.2 Excerpted from Spycher et al., 1999, Section 7

E.7 Anticipated Error Messages

All execution stops built into TOUGHREACT V2.1 are accompanied by a message indicating why the execution was aborted. Other error messages do not lead to a program interruption. Only messages related to the reactive transport part of the program are reviewed below. Error messages originating during heat or fluid flow calculations are the same as for TOUGH2 V.1.11 (Pruess et al. 1996).

E.7.1 From routine: INIT (reads the CHEMICAL.INP file)

Most of these messages are self explanatory and refer to exceeded array dimensions or other error encountered when reading the CHEMICAL.INP file. Array dimension problems can be corrected by reducing the problem size or changing array dimensions in the source file PARAMETE.INC and recompile the program.

Error: maximum number of component species (MPRI) was exceeded. Current max= ____
Execution stop: yes. Self explanatory.

Error: maximum number of minerals (mmin) was exceeded. Current max= ____
Execution stop: yes. Self explanatory.

Error: maximum number of gases (mgas) was exceeded. Current max= ____
Execution stop: yes. Self explanatory.

Error: maximum number of ads. species (mads) was exceeded. Current max= ____
Execution stop: yes. Self explanatory.

Error: MBOUND(parameter of adjustable array) should be great than ____
Execution stop: yes. Too many boundary waters were specified.

Error: MRECH (parameter of adjustable array) should be great than ____
Execution stop: yes. Too many recharge waters were specified.

error in number of primary species in zone= ____ (iwtype)

Execution stop: yes. One of the initial, boundary, or recharge waters does not have data for one of the component species specified in system, or has more components than specified for the simulation. All species listed in the first part of the CHEMICAL.INP file must have a concentration defined for each specified initial, boundary, or recharge water.

error reading chemical control parameters

Execution stop: yes. Error while reading the CHEMICAL.INP file at one of the following locations: initial title; number of initial, boundary, and recharge waters (niwtype,nbwtype,nrwtype); number of sorptions zones (ndtype).

error reading aqueous species of the system

Execution stop: yes. Self explanatory.

error reading minerals of the system

Execution stop: yes. Self explanatory.

error reading gases of the system

Execution stop: yes. Self explanatory.

error reading surface complexes of the system

Execution stop: yes. Self explanatory.

error reading the model of surface complexes

Execution stop: yes. Self explanatory.

error reading initial water zone=___ (iwtype)

Execution stop: yes. Self explanatory.

error reading initial mineral zone= ____ (imtype)

Execution stop: yes. Self explanatory.

error reading initial gas zone= ____ (imtype)

Execution stop: yes. Self explanatory.

error reading initial surface adsorption zone= ____ (idtype)

Execution stop: yes. Self explanatory.

E.7.2 From routine: NRINIT (initial Newton-Raphson iterations)

ERROR (convergence problem in initialization of water composition)

*Please adjust convergence criteria regarding chemical iteration
and initial guess of concentration of primary species*

Execution stop: yes. Self explanatory. This error results in calling routine chdump for troubleshooting (i.e. the last chemistry calculation data are output in the *chdump.dat* file). This error occurs during the initial geochemical speciation of waters specified in CHEMICAL.INP (no minerals, before the first time step). Check the *chdump.dat* file for clues, and also check that

water temperatures specified in CHEMICAL.INP data are not too different than the initial condition of temperature specified in the FLOW.INP file.

E.7.3 From routine: READTHERM (reads the database THERMOK.DAT)

All these messages occur while reading the thermodynamic database and are self explanatory. These indicate improperly-formatted records in the THERMOK.DAT file. All errors result in a program execution stop

Error reading temperature data: stop

Error reading primary species: stop

Error reading secondary species: stop (followed by the species name)

Error reading minerals: stop (followed by the mineral name)

Error reading gases: stop (followed by the gas name)

Error reading adsorbed species: stop (followed by the species name)

Error in opening database file: stop

E.7.4 From routine: READSOLU (reads the file SOLUTE.INP)

There are currently no specific error messages generated while reading the file SOLUTE.INP. The unit number of this file 31. System error messages relating to this I/O number originate while reading this file. Make sure the fixed formats of this file are respected.

E.7.5 From routine: CR_CP (kinetic data calculations)

error in data option for mineral (kinetic)= ____

Execution stop: yes. This message occurs if the flag IDEP for any of the kinetic minerals is not set to either 0 or 1. With this program version, IDEP must always be zero (this flag is specified in the mineral section of the CHEMICAL.INP file).

E.7.6 From routine: NEWTONEQ (Newton-Raphson iterations after 1st time step)

ERROR: chemistry did not converge at node ____ (routine NEWTONEQ)

Species: ____ Error=____ Error limit= ____ relative

Node temperature (C): ____

Program execution was not aborted. Check results!

Execution stop: only if this error occurs at more than fifty grid blocks at any given time step. This error also calls routine chdump for troubleshooting (i.e. the last chemistry calculation data

are output in the *chdump.dat* file). This error occurs during the node-by-node geochemical speciation computations after the first time step (complete system, with minerals and gases if any). Check the *chdump.dat* file for clues on why convergence was not reached. You may need to increase the loop limit (MAXITCH) and/or tolerance (TOLCH) in the SOLUTE.INP file. If boiling occurs, you may try increasing STIMIN or decreasing STIMAX. Chemical convergence may also fail because of errors during transport, resulting in erroneous system compositions which cannot yield a solution to geochemical speciation calculations. In this case, the time step may be decreased and/or the Courant number option enabled (ICOUR=1). Depending on the type of problem, chemical speciation in closely spaced grid blocks can be skipped by setting DIMAX > 0 (last resort).

Error: Negative concentration for species _____

Execution stop: no. Self explanatory. A concentration may temporarily become negative during the chemical Newton-Raphson iterations, but should not remain negative. This error may indicate problems to come. It is rarely encountered.

E.7.7 From routine: LUDCMP (linear equation solver)

This routine is called during the Newton-Raphson iterations to compute geochemical speciation.

Singular Matrix in Chemical Solver, STOP

Execution stop: yes. Self explanatory. This indicates an ill-defined chemical system. This error results in calling chdump to output the last geochemical speciation data in the *chdump.dat* file. A phase-rule violation or inclusion of minerals (at equilibrium) with identical stoichiometries in the simulation will cause this error. In some cases, divergence and over/underflows during Newton-Raphson iterations (sometimes related to transport problems) may cause this error even though a true singularity has not occurred. Check the *chdump.dat* file for clues on why the error happened.

Appendix F: Input/Output Changes from V2.1

A general description of TOUGHREACT V2.1 input and output files and formats is provided in Appendix D. Only changes from V2.1 (see Spycher et al., 1999) are described here.

FORTTRAN format notations are to be used to describe fixed formats (A for strings, I for integers, F for reals, X for spaces, etc.). Free formats are to be described using a modified FORTTRAN notation, with format letters specified without a length. Free formats for arrays may be preceded by a variable (e.g. `ntmp*F`) which indicates the maximum dimension of the array. These dimensions are generally to be defined in the `PARAMETER.INC` source file. Character strings in free format must be enclosed in single-quotes. As a reminder, free format string variables below are listed with names between quotes.

F.1 FLOW.INP input file

This file is to be in standard TOUGH2 input format (Pruess 1991) and to be consistent with EOS3 V1.4 input formats (Wu et al. 1996 and 1999), except for different assigned ISOT values, as noted below.

The following new flag values shall be implemented in the ROCKS input block:

- | | |
|----------|--|
| ICP = 10 | Flag for capillary pressure function (see TOUGH2 user's guide): same function as with ICP = 7 (van Genuchten model), but the capillary pressure is linearly interpolated at liquid saturations below $S_{\text{residual}} + (S_{\text{residual}} \times \text{factor})$, using the slope of the capillary pressure-saturation curve at that point. S_{residual} is entered as CP(2) (as in V2.1) and <i>factor</i> is entered as CP(4) (in place of P_{max} in use if ICP = 7). |
| CP(6) | Active-fracture model parameter γ as defined in equation A-4 (only effective if ICP = 7 or ICP = 10). Leave blank or zero to disable the active-fracture model, and specify this parameter only for fracture rock types. |

The following new flag values shall be implemented in the CONNE input block:

- | | |
|------------|--|
| ISOT = -10 | Enables the active-fracture model (in conjunction with non-zero CP(6) values) for the specified connection. Set ISOT = -10 only for fracture-matrix connections. |
| ISOT = -15 | Enables downstream weighting of permeabilities at the specified connection (Note!!!: the same option in EOS3 V1.4 is enabled with ISOT = -2) |

The following new flag values shall be implemented in the PARAM input block:

- | | |
|-------------|---|
| MOP(21) = 5 | New solver (for flow computations): conjugate gradient stabilized (preconditioned) with LU factorization. |
|-------------|---|

A new flag (*ktab*) shall be added to the first record of the GENER input block to implement time-varying thermal conductivities:

- 1st record of GENER block (A3,I2,A3,I2,4I5,5X,A4,A1,3E10.4,I2): EL, NE, SL, NS, NSEQ, NADD, NADS, LTAB, TYPE, ITAB, GX, EX, HG, *ktab*

EL, NE, SL, NS, NSEQ, NADD, NADS, LTAB, TYPE, ITAB, GX, EX, and HG are as previously defined for V2.1 (see TOUGH2 users' guide, Pruess 1991).

ktab If non-zero or non-blank, *ktab* enables input of thermal conductivity factors into the GENER block. With this option, LTAB must be blank or zero and TYPE must be set to "HEAT". *ktab* must be set equal to the number of time-variable conductivity factors that need to be input. Time values in the GENER block are then input as standard TOUGH2 generation times (F1(L), L = 1, *ktab*), and corresponding thermal conductivity factors are input as standard TOUGH2 generation rates (F2(L), L = 1, *ktab*). With this option, the thermal conductivity (input in the ROCKS block) is multiplied by the factors F2 at corresponding times F1.

F.2 SOLUTE.INP input file

Input formats are to be the same as for V2.1, but a record is to be added to the file. This file is therefore not backward compatible with V2.1 without some minor editing.

Records 1 to 10 are the same as in V2.1. A new record is then to be inserted as follows:

- 11th record (A20,A60): OUTGAS, TITLE
Name of file for output of gas plot data (TEC_GAS.DAT) and any title

Next records are the same as in V2.1 (but offset by one record) until record 18 (corresponding to record 17 in V2.1), where a new flag (MINFLAG) is to be added:

- 18th record (7I5): NWTI, NWNOD, NWCOM, NWMIN, IWCOMT, MINFLAG

NWTI, NWNOD, NWCOM, NWMIN, IWCOMT are to be as previously defined for V2.1. MINFLAG is to be a new flag to indicate output units for gas and mineral amounts: MINFLAG = 0 or blank, no change from V2.1 (output in bars for gases and moles/m³_{medium} for minerals); MINFLAG = 1, gas and mineral amounts are to be output in volume fractions (of the medium).

A new solver is to be available by setting ISOLVC, in the third record of this file, as follows:

ISOLVC = 5 New solver (for transport computations): conjugate gradient stabilized (preconditioned) with inprilete LU factorization.

A new option for gas diffusion coefficients is enabled by setting DIFUNG, in the 13th record of this file, as follows:

DFFUNG = -1.0 Enables calculation of the gas diffusion coefficients as a function of temperature and pressure using Equation A-1.
DFFUNG > 0 In this case DFFUNG is the gas diffusion coefficient (not multiplied by tortuosity, as was the case in V2.1).

F.3 CHEMICAL.INP input file

Input formats are to be the same as for V2.1, but a few flags are to be added to the file. This file is therefore not backward compatible with V2.1 without some minor editing.

All record formats and variables are to be the same as in V2.1, except for the mineral records and initial mineral zones records.

In the mineral records, a flag (NPLAW) is to be added to specify the precipitation rate law as follows:

- 3rd mineral record, only if IKIN > 0 and IDSPRE = 2 or 3 (F,I,7F,I): RKPREC, IDEPREC, CK1PREC, CK2PREC, EAPREC, ACFPREC, BCFPREC, CCFPREC, RNUCL, NPLAW

RKPREC, IDEPREC, CK1PREC, CK2PREC, EAPREC, ACFPREC, BCFPREC, CCFPREC, and RNUCL are to be as previously defined for V2.1. Specify NPLAW = 0 (cannot be left blank) for the default rate law (same as in V2.1), and NPLAW = 1 for the new rate law (equation A-7).

In the initial mineral zones records, a flag (IMFLG) is to be added to specify the units of input surface area as follows:

- 2nd mineral record (in mineral zones), only if IKIN4 = 1 (2F,I): RAD, AMIN, IMFLG

RAD is as previously defined for V2.1. AMIN is the mineral surface area in [$\text{cm}^2/\text{g}_{\text{mineral}}$] if IMFLG = 0 (cannot be left blank), and in [$\text{m}^2/\text{m}^3_{\text{mineral}}$] if IMFLG = 1.

F.4 THERMOK.DAT input file (thermodynamic database)

Input formats are to be the same as for V2.1, but a few variables were added/changed in the gas records. This file is therefore not backward compatible with V2.1 without some minor editing.

All record formats and variables, except for the first record of each gas entry and the last record of the gas block, are to be the same as for V2.1. Two variables (DMOLWT and DMDIAM) are to be added to the third gas records as follows:

- 1st record (A,2F,I,mpri*(F,A)): NAME', DMOLWT, DMDIAM, NCPG, (STQG(i), NAM(i)', i=1,NCPG)

NAME, NCPG, STQG, NAM, and NCPG are to be as defined for V2.1. DMOLWT is the gas molecular weight in [kg/mol] and DMDIAM is the gas molecular diameter in [m]. These two variables are to be used to compute the gas diffusion coefficient according to equation A-1 if DFFUNG (the gas diffusion coefficient variable) in the SOLUTE.INP file is set to -1.0.

The maximum total number of gases allowed in the program is to be *mgas* (defined in source file PARAMETE.INC). After the data for all gases have been listed (three records per gas), the last record to end the gas data must be as follows:

F.5 Output files

Output files are to be the same as in V2.1, except that the TEC_MIN.DAT file of V2.1 (which then included gas data) is to be split into two files with V2.2: TEC_MIN.DAT (without gas data) and TEC_GAS.DAT for the gas data. Also, compared to V2.1, additional variables are to be output in plot files as follows: porosity and permeability (m^2) in TEC_MIN.DAT and total pressure (Pa) in TEC_GAS.DAT. Mineral and gas amounts in these files are to be output in units that are set by the value of MINFLAG (see Section F.2).

F.6 References

- Pruess K., *TOUGH2-A General Purpose Numerical Simulator for Multiphase Fluid and Heat Flow*. Report LBL-29400, UC-251, Berkeley, California: Lawrence Berkeley National Laboratory, 1991. NNA.19940202.0088.
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