

SHPECK - A Geochemical Speciation Modeling Software

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Abstract

Geochemical speciation modeling software are used for calculating the distribution of dissolved solute and complex compound species between solutes and aqueous complexes, and also to compute saturation indices of minerals. In this work we introduce *SHPECK*, a software program developed to model geochemical speciation using the mass-balance conditions and the phase rule [Garrels and Christ 1965]. *SHPECK* can be used to model the chemical equilibrium interaction between elements and solutes. It provides an interactive and intuitive user interface, as well as the support of a relational database structure that handles the management of thermodynamic and material properties data used for the geochemical modeling. We review the available geochemical modeling software (only programs that provide speciation modeling), highlighting relevant characteristics as: input and output options; user interaction; file formats; and software environment and installation procedure. The software *SHPECK* is validated by modeling the diagenetic reactions observed in a siliciclastic geological system, and by performing a comparative study with other modeling software package. In addition to this, a database comparison was addressed and the results demonstrate a substantial improvement on the performance by the use of the *SHPECK*'s relational database comparing to the existing approaches.

Keywords: Geochemical modeling, Chemical Equilibrium, Geochemistry, Multiphase System, Law of Mass-Action, Nonlinear simulation, Software Engineering, Computational simulation,

1. Introduction

Geochemical speciation modeling describes the process of simulating chemical reactions that occur between the water (solvent) and chemical elements. The motivation of this

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study is to provide a better understanding of the diagenesis process of sediments during the burial process, especially the water-mineral chemical interaction. Applications of geochemical models are essential in several environmental problems, such as calculating compositions of natural ground or surface waters, and precipitation and/or dissolution of minerals. A geochemical speciation modeling software is responsible for calculating the distribution of dissolved species into solutes and aqueous complexes, and also for computing saturation indexes for different mineral.

In this work, we present a software system, named as *SHPECK*, which allows the simulation of the chemical equilibrium reactions occurring in a geochemical system using the mass-balance approach and the phase rule as described by Garrels and Christ [Garrels and Christ 1965]. The software system receives as input any general combination of chemical elements, species, and reactions, allowing the user to create several different environments, simulations and, therefore, and to fully control any aspect and configuration of the model. Moreover, our work shows a thorough analysis of the available existing solutions, making clear the uniqueness of the proposed computational approach for the geochemical modeling problem.

The software *SHPECK* also provides an interactive user interface that allows the users to more conveniently setup the simulations. The user interface used a relational database that further improves database archiving and managing, which is an option absent in any of the other alternatives. The flexibility of parameter configuration offered by the visual interface further extends the possibilities of experimentation with the software, and this is a unique characteristic among commercial software products for geochemical speciation. Limitations on the database structure and input capability strongly restrict the potential use of the the alternative geochemical packages. Therefore, *SHPECK* is unique among the geochemical speciation models by implementing modern computing technology, absent in other similarly tasked software, and also employing efficient numerical methods.

2. Review of Available Geochemical modeling Software

Geochemical modeling software have been in use since the 70s to solve: speciation; determination of mineral saturation index; adjustment of equilibrium for minerals; mixing of compositionally distinct waters; calculation of stoichiometric reactions; mixing of solids, fluid and gas phases; calculation of equilibrium and kinetic reactions between solutes and minerals; and reactive transport. While a large number of computational models have been developed over the years, a number of them are particularly well known and available to general public. Thus, three models representing the large number of similar software have been identified and compared in this study: *EQ3/6*, *PHREEQC* and *MINTEQ*. We had given special attention to the geochemical modeling approaches and the software engineering aspects adopted in their implementations, providing the comparison when the information is available.

EQ3/6 consists of two programs: *EQ3* is a speciation code whose results *EQ6* subsequently processes. It is a software package for geochemical modeling of aqueous systems written in *FORTRAN77*. *EQ3/6* includes a speciation-solubility solver, which is useful for

analyzing groundwater chemistry data, calculating solubility limits and determining whether certain reactions are in partial equilibrium or in state of disequilibrium. It also offers a reaction path calculation that models water/rock interaction or fluid mixtures. EQ3/6 supports several thermodynamic data files. These data files also include supports for *Davies*, *B-dot*, *Debye Huckel* equations used to approximate activity of solutes in water, as well as data for standard-state properties of solutes. Currently available versions run under *UNIX-like* environments, and the full package distribution requires a paid license. Additional details of the software can be found in [Wolery 1979], [Wolery et al. 1990] and [Wolery 1992].

PHREEQC stands for *PH* *RE*dox *E*quilibrium in *C* language, and is largely used geochemical modeling software obtainable from the *USGS*. Versions for Windows and UNIX-like are available. A detailed description of the package can be found in [Parkhurst 1995]. It was designed to perform a wide variety of low-temperature aqueous geochemical calculations based on an ion association aqueous model. It includes functionality for:

- Speciation and Saturation Index calculations;
- Batch reaction and one-dimensional (1D) transport calculations involving reversible reactions (including aqueous, minerals, gas, solid-solution, surface-complexation, and ion-exchange equilibrium) and irreversible reactions (including specified mole transfer of reactants, kinetically controlled reactions, mixing of solutions and temperature changes);
- Inverse modeling, which finds sets of mineral and gas mole transfers that takes into account differences in water composition.

MINTEQ [Felmy et al. 1983] is a geochemical program for modeling aqueous solutions and their interactions with hypothesized assemblages of solid phases. It has a particular inclination to calculate equilibrium composition of dilute aqueous solutions. The model is useful for calculating the equilibrium mass distribution among dissolved species, adsorbed species and multiple solid phases, limited by a simple treatment of the reactions. It was originally developed in *FORTRAN77* by the *Battelle Pacific Northwest Laboratories (BPNL)* and continues to be maintained by the *Environmental Protection Agency (EPA)* to perform the necessary calculations regarding waste, sediments and ground water interaction. *MINTEQ* does not consider the kinetic reactions and works at fixed temperature (25 degrees Celcius). An extensive database is included in the software [Brown and Allison 1987] [Allison J. D. and Novo-Gradac 1991]. The latest update on *MINTEQ* dates from 1990. Limited improvements on the usability and calculations have been added since, and the new version is named *MINTEQA2*. The new version supports the thermodynamic database from the *USGS*. In this article only the *MINTEQA2* is reviewed.

2.1. Input and Output Format

EQ3/6 and *MINTEQA2* work with input files, more than one, that are generated manually and given to the program as arguments. It is important to note that they must be mutually consistent with the options and methods. Wrong chemical data or inconsistencies

in the chosen algorithm will lead to erroneous results. *PHREEQC* works with the same approach, however a visual plugin [Apello 2011] and a *GUI* [Post 2011] make it possible to reduce the error. It is often very tedious and time consuming to generate input files for simulations. Output formats on the three software systems are similar: text files with blocks of information reporting input options, variables and results of the simulations.

2.2. User Interaction

EQ3/6, *MINTEQA2* and *PHREEQC* (*UNIX-like* distribution) develop all the interaction with users through the command line, since there is no graphical interaction. Once the simulation process starts, any change in the parameters requires to kill the process manually through the operating system and run the simulator again. The *PHREEQC* package provides a version for Windows [Post 2011], however it has not been updated since 2011. This version allows the user to generate input files, however there are no error verification or analysis capabilities.

2.3. File formats

All three software mentioned work with text input files (*ASCII*), which can be created and edited using any text editor.

2.4. Thermodynamic database

The three software use flat files as thermodynamic database. This mechanism has demonstrated along the years several potential issues: duplication of the information; non-unique records; difficulty updating and maintaining; inherently inefficient; rigid (difficult data format); and insecure [?].

2.5. Software environment and installation procedures

EQ3/6 has no self-extracting installer, and the installation process requires some level of proficiency with command prompts and *DOS*. Furthermore, *EQ3/6* is available only for *Windows*; the *UNIX* distribution is no longer available. *PHREEQC* and *MINTEQA2* have *Windows* installers available for download. *MINTEQA2* distribution on *UNIX* environment consists of the source code, thus requiring the user to compile and link. *EQ3/6* and *MINTEQA2* were written in *FORTRAN77* while *PHREEQC* is written in *C*.

3. SHPECK

SHPECK is a geochemical speciation modeling software capable of calculating the distribution of dissolved species between free ions and aqueous complexes and also the saturation indexes for different mineral phases. *SHPECK* was developed using *C++* and is modelled following the concept of *Model-View-Controller (MVC)* [Gamma et al. 1994]. The architecture is presented in figure 1.

MVC is an architectural pattern that divides our software into three interconnected parts:

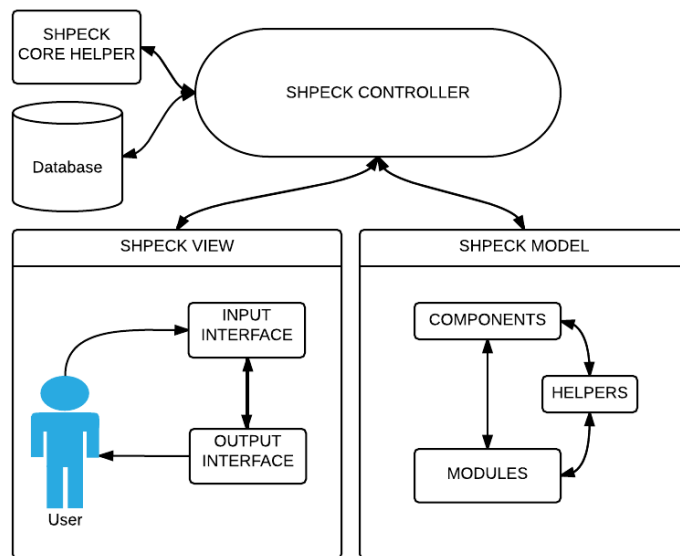


Figure 1: Architecture of the *SHPECK* software

- Model: It represents, for example, the algorithm and math behind calculating the activity coefficient by Debye-Hueckels' formula.
- View: It is a form of visualization of the state of the model. For example, the solutes that the user wants to add into the chemical composition of the water.
- Controller: It offers ways to change the model's state. For example, define how to calculate the activity coefficient according to the user's choice or the value of the ionic strength.

3.1. Graphical User Interface

In existing geochemical modeling software systems, the *GUI* are either poorly implemented, or not implemented at all. The *GUI* of *SHPECK* enables a user to interactively construct a geochemical system for modeling. *SHPECK*'s *GUI* uses tabs to address different criteria and input types, as it is presented in detail below:

- Configurations panel (figure 2): this panel allows users to view and manipulate basic system settings and to control temperature, activity coefficient calculation method, the number of iterations, solver options, and other numerical method parameters.
- Chemical phases in the water panel (figure 3): this panel allows users to create and edit composition of waters that are used in the model. This section has access to the complete catalog of solutes database. The solutes in the tab with non-zero concentrations are included in the simulations.

- Results panel (figure 4): This panel allows users to examine input information and simulation results, such as temperature, ionic strength, pH of the solution, final concentration for the solutes, saturation indices, etc.

Figure 2: Configuration tab

3.2. Database

A relational database is used in *SHPECK* to bypass the problems associated with the use of text files used as databases in other software, and to provide flexible and rapid, visually effective correlation between information required and available through the database enquiries. In addition, relational databases offer three important advantages:

- Memory saving, since it does not require the complete database to be fully load in main memory during the run time. The program fetches the information only when required by each module.

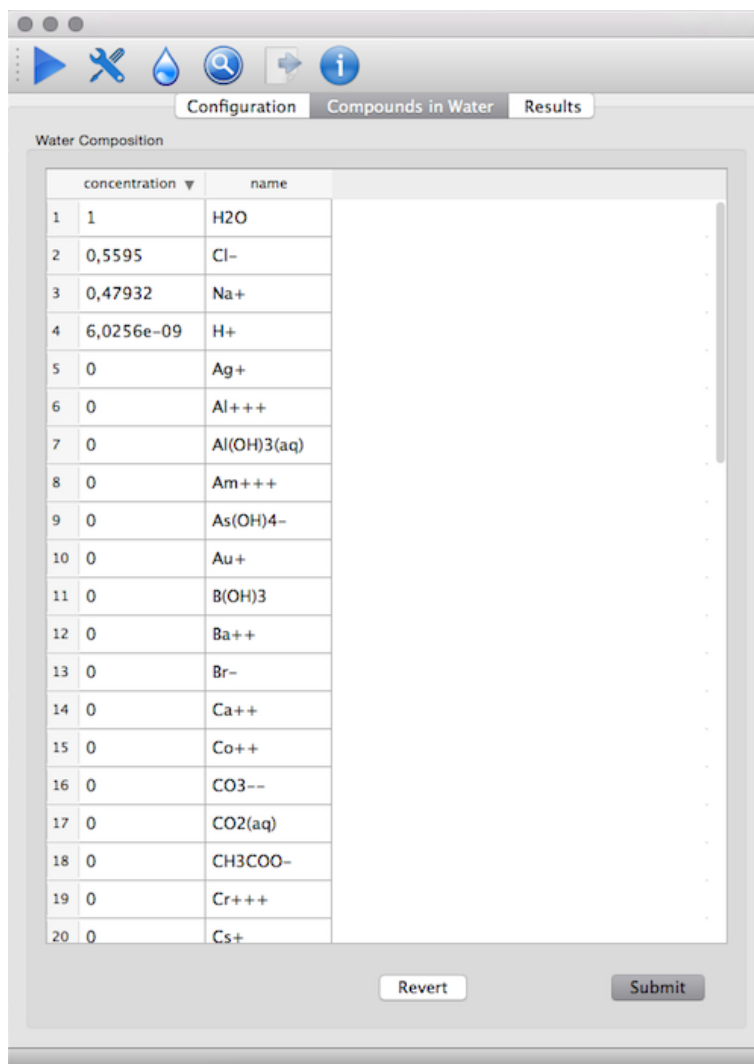


Figure 3: Water composition tab

- Direct access to relevant information: SQL queries are used to fetch only the relevant information, whereas sequential search is required when text files are used. Database queries and concatenation of queries result in a faster and more efficient use of the available resources.

The database model, presented in details in figure 5, was defined after studying the algorithm and determining what would be the structure and the information needed. The relational database embraces all the important data required for a geochemical modeling software, and the structure was defined to both organize and compact the structure, making the information access more efficient.

We adopted a *SQLite* database [Hipp 2015], which is a software-library that implements a self-contained, transactional *SQL* open source database engine. Currently it is the most widely deployed *SQL* database engine in the world.

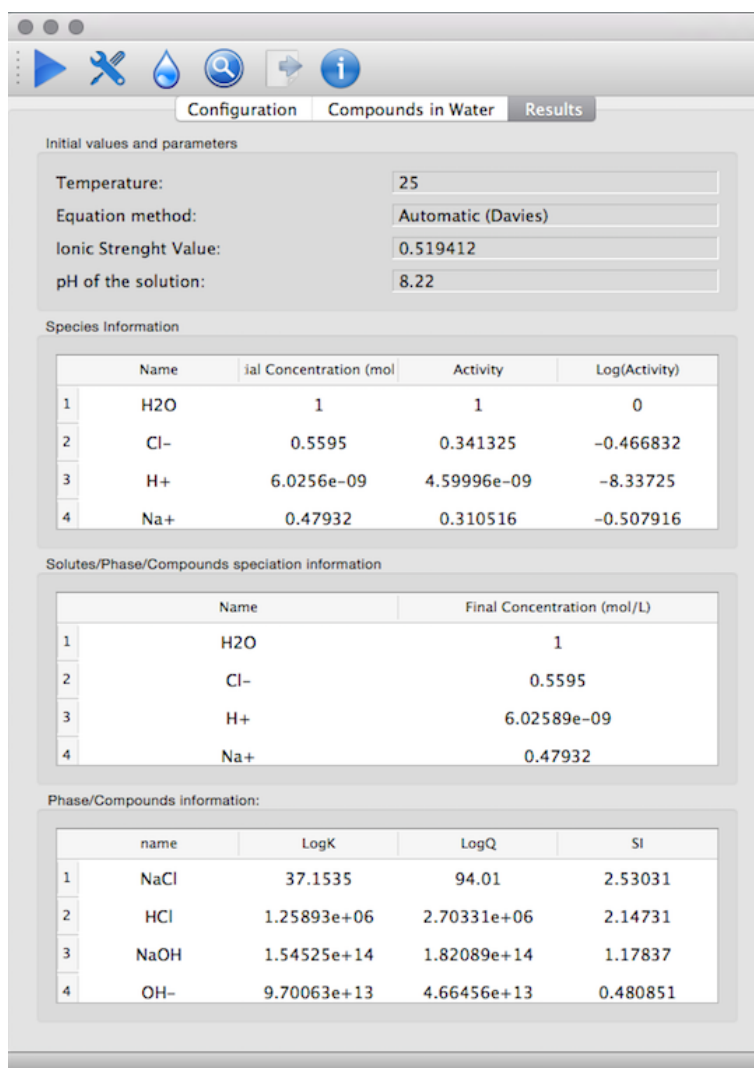


Figure 4: Results tab of *SHPECK*

SHPECK's database is composed by the information about elements, solute species, compounds, reactions and thermodynamic constants from the *Lawrence Livermore National Laboratory (LLNL)* thermodynamic dataset. A parser for *LLNL* flat file database was created to extract this information from text files and to populate the database tables. The same text files are used as database in *EQ3/6*, *PHREEQC* and other geochemical codes.

Once the structure and the technology of our database were defined, a program was implemented to extract (i.e., parse) the information from the flat file database. The parser was generated in order to ensure that all of the data and delimiters were identified and treated properly, because of irregularities found in the flat files (i.e. wrong tab or space forming, different encoding of the files, etc).

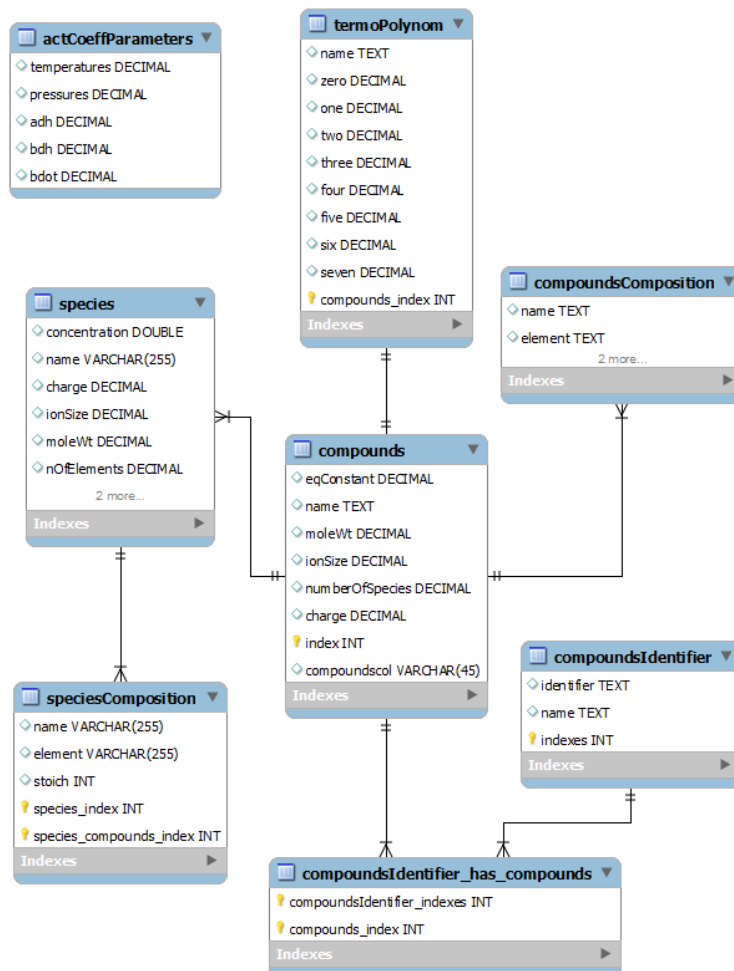


Figure 5: ER Diagram of the database

4. Verification and Validation

To test and validate the simulator, we model the diagenetic reactions observed in Snorre Field reservoir sandstones of Norwegian North Sea. The main reservoir horizons of the field are the fluvial sandstones in the upper member of the Upper Triassic Lunde Formation and the Upper Triassic to Lower Jurassic Statfjord Formation [Hollander]. The sandstones sampled for this study, according to Morad [Morad. S. 1990], belong to the upper member of the Lunde Formation. The sandstones are dominantly fine- to medium-grained and arkosic with framework constituents of quartz (40-80%), K-feldspar (5-12%), plagioclase (15-45%), muscovite, biotite and clay minerals that include smectite, mixed-layer clay minerals, chlorite and subordinate amounts of kaolinite and illite. Subordinate rock fragments include intraformational mudstone and carbonate clasts and extrabasinal grains of quartz-feldspar-mica aggregates that probably represent granitic rocks and/or schist or gneisses. Mica and detrital clay minerals seldom make up more than 2% of the total mineral content in the

sandstones. Diagenetic clay minerals include pore-filling kaolinite and pore-lining smectite, mixed-layer chlorite-smectite, and chlorite. Other cements include the carbonates (0.0-25%) which play a significant role in porosity reduction in some of the sandstones. Authigenic overgrowths are primarily quartz, anatase and minor albite, pyrite and barite.

The descriptions presented in [Morad. S. 1990] of the diagenetic reactions that take place in the Snorre Field allowed us to generate results to carry out a computationally comparative study. Morad's work describes the texture, origin, chemistry of the sandstones reservoirs in terms of the water composition and temperature. We model and compare the same diagenetic environment using *SHPECK*, *PHREEQC* and *MINTEQA2*. The water composition is detailed in [Nordstrom 1979].

As stated in [Morad. S. 1990], the model presented in [Egeberg and Aagaard 1988] calculates activities of the various ions of formation waters using the ion association model (originally described in [Wigley 1977]). The thermodynamics data used are given in [Helgeson and Kirkham 1974], [Helgeson and Kirkham 1976], [Walter and Helgeson 1977], [Helgeson H. C. and Flowers 1981].

The activity diagram generated for a known temperature and log activity ratio of Potassium to Sodium ions of [Aagaard 1990] is provided as input to *SHPECK* model (Figure 6). The results show a consistent pattern: as the temperature rises the potassium activity becomes higher than sodiums, which means that the phases associated to the ion potassium (i.e. K-feldspar, Illite, etc) are dissolving.

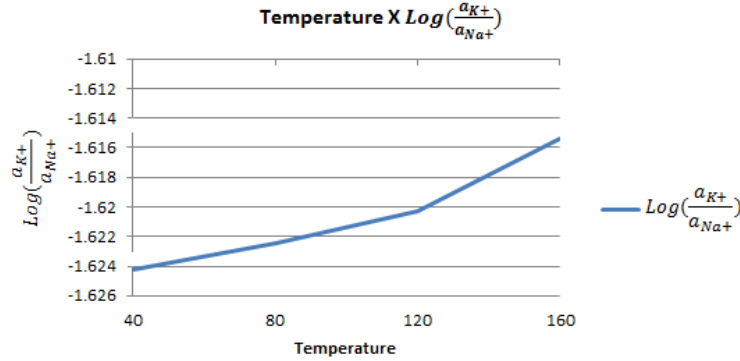


Figure 6: Log activity ratio of Potassium to Sodium ions using the results from *SHPECK*

4.1. Comparative study

By modeling the same environment using three distinct software packages, we attain a relevant comparison among the numerical methods and algorithms.

Chemical composition of the water adopted in the models is taken from [Nordstrom 1979] which provides the chemical composition of the seawater (table 1). The comparative study tested temperatures from 25°C to 100°C. In *MINTEQA2*, due to limitations of its thermodynamics equilibrium database, the maximum temperature available is 100°C.

Table 1: Chemical composition of the solution in the sea water at 25° in mM/LC									
Al ³⁺	K ⁺	Na ⁺	Ca ²⁺	Mg ²⁺	Fe ²⁺	SiO ₂	SO ₄ ²⁻	Cl ⁻	pH
7.59e-5	10.45	479.32	10.53	54.39	3.66e-5	0.073	28.893	559.5	8.22

We aim to model the diagenetic processes that best represent the behavior of ions in the water-rock interactions. Figures 7, 8, 9 and 10 present the most representative ions of the solution.

The results of *SHPECK* are similar to those from both *PHREEQC* and *MINTEQA2* in most of the cases, especially for temperatures under 100°C. Discrepancies occur in *SHPECK* and *PHREEQC* results when temperatures are higher than 100°C. This discrepancy is due to incomplete equilibrium constant K defined in the *MINTEQA2* database.

This is a known issue from *LLNL* thermodynamic dataset: sometimes equilibrium constants have no measures in literature and are treated as unknown. *SHPECK* adopts the nearest equilibrium constant known value. Unfortunately we do not have access to the whole softwares details in order to describe how they treat this issue.

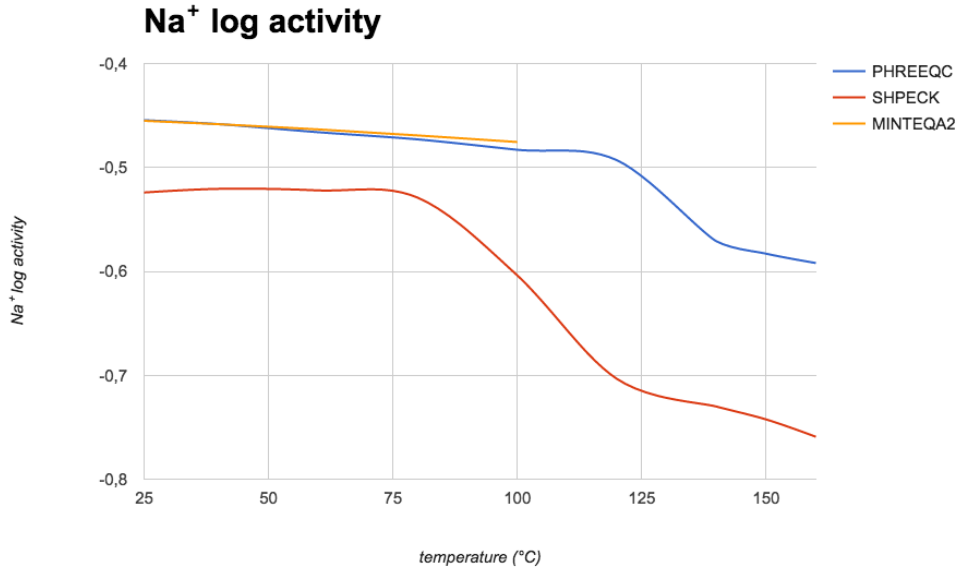


Figure 7: Na⁺ log activity comparative study

4.2. Database Evaluation

All geochemical models used for comparison studies use text files as databases. The goal of this section is to make clear the difference and, more importantly, the benefits of *SHPECK*'s relational database. Most of the information inside a geochemical database is related to each other (i.e. a mineral phase is described by a reaction, a reaction is composed by solute species, and a solute species is composed of chemical elements). Relational

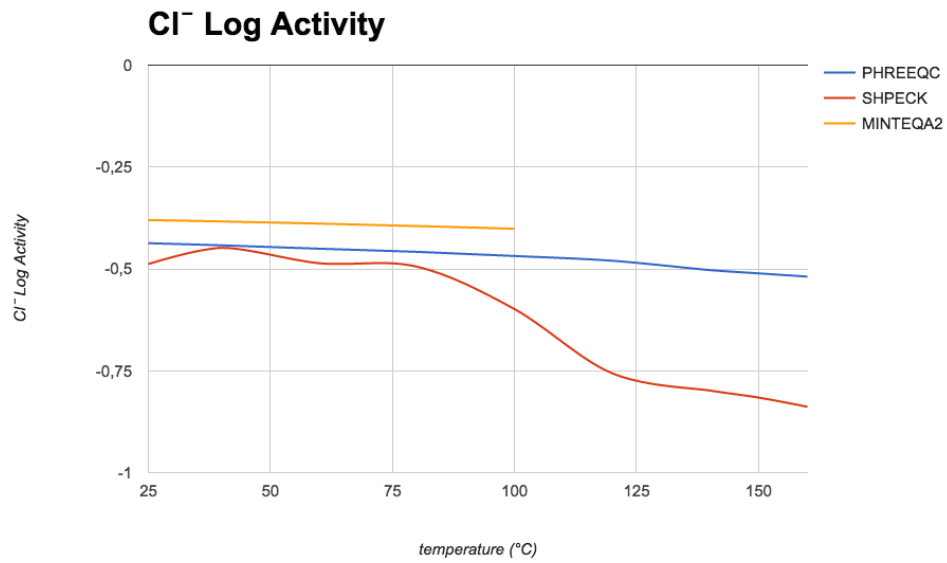


Figure 8: Cl⁻ log activity comparative study

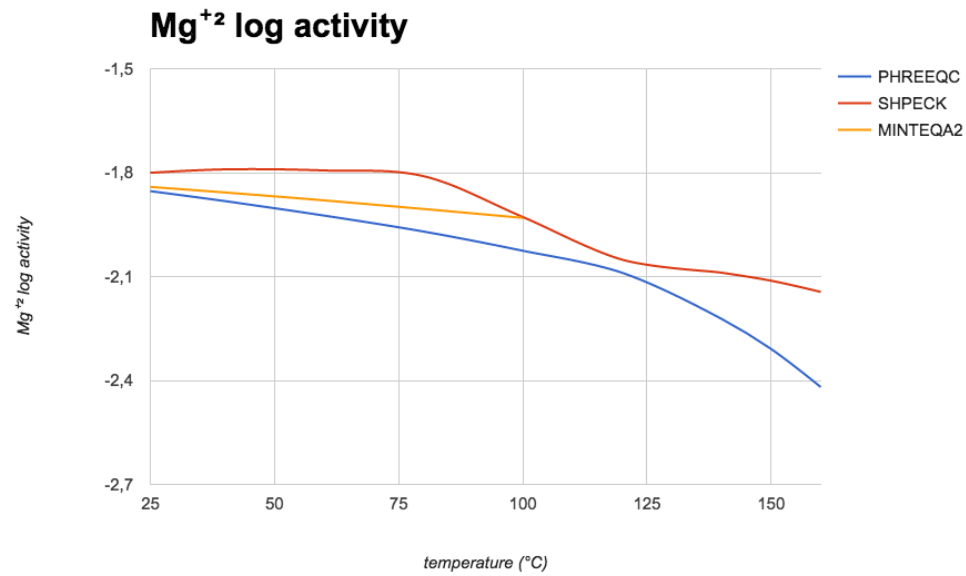


Figure 9: Mg⁺² log activity comparative study

databases sophisticated built-in query and sorting statements make the organization of the data convenient, and this significantly improves the performance and robustness of the application. On *SQLite* databases, the data can be accessed using *SQL* queries that reduce

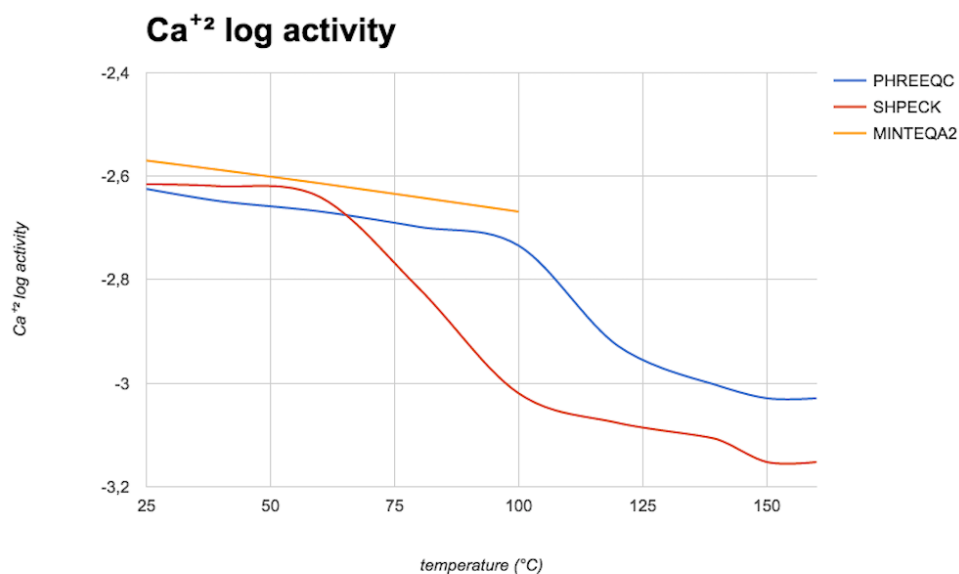


Figure 10: Ca²⁺ log activity comparative study

the complexity and increase the speed on information retrieval. Code 1 shows a typical query issued from *SHPECK*'s *SQLite* query. The options and the flexibility afforded by the use of relational database cannot be compared to the options available when textfile-based databases are used.

Code 1: *SHPECK*'s *SQLite* example query

```
SELECT compoundsComposition.element, compoundsComposition.name,
    compoundsComposition.stoich, compounds.eqConstant, termoPolynom.zero,
    termoPolynom.one, termoPolynom.two, termoPolynom.three, termoPolynom
    .four, termoPolynom.five, termoPolynom.six, termoPolynom.seven FROM
    termoPolynom, compoundsComposition, compounds WHERE
    compoundsComposition.name = compounds.name AND termoPolynom.name =
    compounds.name AND compounds.name = XXXXX;
```

Important to note that at each simulation the value XXXX in Code 1 is updated to the simulations solute name. It is also interesting to point out that the information fetched with this query comes from three different tables, and thus, many relevant information for the simulation is fetched at once.

4.3. Time analysis

The response time is considered as the sum of the processing time and the time waiting for the availability of the resource. It is necessary to understand that until the software has received all the information requested from the database it is inactive and in a standby mode. To analyze the response time within a geochemical analysis point of view, we discuss not only the access time, but also the implications of that information.

When fetching any information from a thermodynamic database, it is important to take

into consideration additional allied data will also have to be retrieved. For example, when fetching a reaction data, the basic information consists of the compounds that take part in this reaction and the related stoichiometric values. Behind this action, the database must also provide information about the compounds (i.e. charge, ion size, molecular weight, elements in that species, formula, and molar volume), as well as the reaction (i.e., thermodynamic equilibrium constant coefficients, etc).

Figure 11 indicates the time elapsed (in seconds) that it takes to retrieve all necessary information related to a chemical reaction from the database. In this example, we simulated from 20 to 580 randomly selected reactions in the database. It is possible to see that *SHPECK*'s database has improved approximately 40% in the average time elapsed to fetch the information when compared to the procedure used by other models.

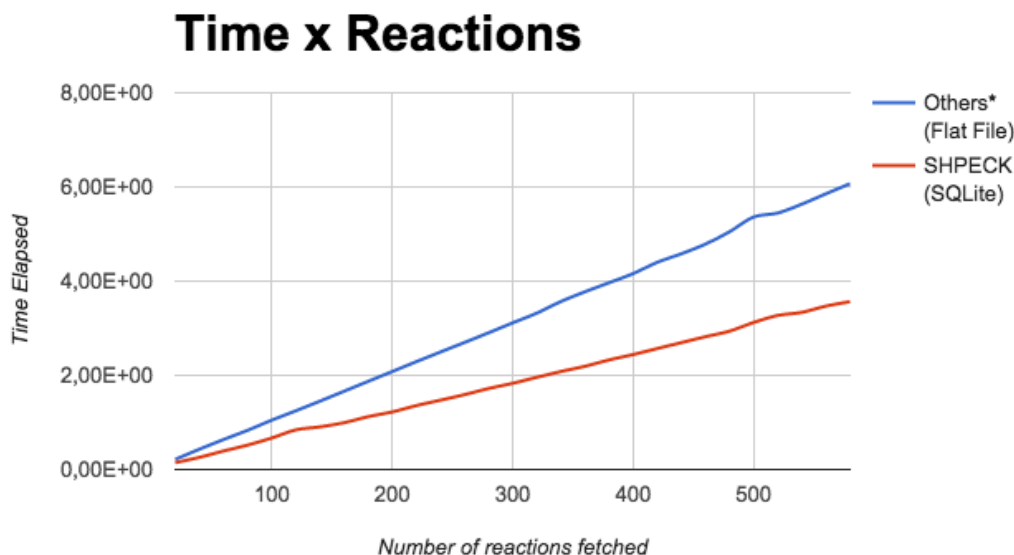


Figure 11: Time elapsed in seconds X Reactions Accessed

5. Concluding remarks

This work provides details of *SHPECK*, a modeling software package intended to raise the quality of available geochemical speciation. The program innovates by including a visual user interface that allows to parametrize the reaction simulation, as well as a relational database architecture component. The user interface implements a dynamic tool that facilitates the simulation case preparation. *SHPECK* accepts any general combination of chemical elements, mineral species and chemical reactions, along with definitions of boundary contour conditions of temperature, pH, pressure, etc.

SHPECK calculates the distribution of dissolved solutes in the water and saturation indices of various mineral phases. *SHPECK* calculates the speciation of a solution based

on a set of mass-balance equations, which are solved iteratively using the Newton-Raphson numerical method.

The software was tested and its results compared with those obtained by modeling the same study case with other geochemical modeling software. The results produced by *SHPECK* were found to be consistent with those of the alternative models.

SHPECK's results accuracy is equivalent and inside the expect range to the study case developed in this work. The results show that all the three software compared have nearly the same results with the same inputs. Slight differences are found when simulating with temperatures higher than 100°C, which appear to arise from discrepancies in the thermodynamic properties of the equilibrium constant.

We plan to improve *SHPECK* by adding kinetic reactions to the processing core (and this is a delicate topic since adding kinetic reactions implies not only working with mass balance equations but also with chemical elemental mass evolution). The elemental mass evolution describes mass change through mass transfer and kinetic reactions of solids and solute-solute interaction - described in details in [Park 2014]. Moreover, a distribution platform where users can register, download *SHPECK*, interact, and collaborate in forums is planned for near future.

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