CROSSING SCALES WITH COMPUTATIONAL TOOLS:
APPLICATIONS TO DIVALENT SILICATE DISSOLUTION

A Dissertation in
Chemistry
by
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ABSTRACT

Computational tools are used to cross spatial scales during investigations of geochemical reactions and mineral surfaces. Density functional theory (DFT) calculations are employed to investigate M–O (M = Mg$^{2+}$, Ca$^{2+}$, and Ni$^{2+}$) bond breaking and H$_2$O exchange using a H$_2$O molecule and molecular sized clusters analogous to sites on silicate mineral surfaces. The barrier heights for hydrolysis of protonated, neutral, and deprotonated Mg–O–Si sites on the forsterite surface were determined. These barrier heights were used to calculate the rate constants, and in turn, a rate for the release of Mg$^{2+}$ due to the breaking of the Mg–O bond. In a second set of calculations, hydrolysis of protonated M–O–Si (M = Mg$^{2+}$, Ca$^{2+}$, and Ni$^{2+}$) sites was investigated to determine whether H$_2$O exchange or bond breaking occurred for Ni$^{2+}$, Mg$^{2+}$, and Ca$^{2+}$ silicate molecular clusters. Here again, the barrier heights are used to calculate rate constants for release of these metals from protonated sites on silicate surfaces. A comparison with experimental data is given, and experimental trends are replicated.

Density functional theory molecular dynamics (DFT-MD) simulations enable the use of a unit cell sized system and allow for an investigation of several reaction sites on the mineral surface. The forsterite (100) and (010) surfaces were investigated to determine the most stable structures for these surfaces when initially covered with all H$_2$O molecules or OH groups. The surfaces yielded similar structures comprised of H$_2$O, OH, O$^-$, and O$_{br}$ sites, and a true forsterite surface likely has a distribution of all of these sites. These surfaces were simulated in the presence of bulk water to investigate the surface structure at the aqueous-mineral interface over time, and H$^+$ and H$_2$O transfers between groups throughout the surface and between the solution and the surface were observed.
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Chapter 1

Introduction

1.1 Increasing Levels and Storage of Atmospheric CO$_2$

The rising levels of CO$_2$ in the atmosphere$^{1,2}$ have increased our need for a better understanding of the global carbon cycle. The global C cycle describes the transformation of C substances throughout the Earth between the atmosphere, bodies of water, Earth’s surface, and biota.$^3$ The interplay between anthropogenic CO$_2$ and C mitigation by minerals has recently received much attention$^{4-10}$ due to the potential to store C long-term. These studies provide insight into how geological systems play a role in the C cycle.

The ability of a particular mineral to mitigate C depends, in part, on the release of metal ions from its structure. In addition to dissolution of minerals for CO$_2$ storage, this process occurs on Earth’s surface and affects groundwater chemistry, erosion, and the concentrations of soil nutrients.$^{11}$ Dissolution of minerals controls numerous life processes on Earth, and particular focus has been given to investigations of silicate minerals because they are the most abundant class of minerals on Earth’s surface.$^{12}$

Silicate minerals are desirable choices for geologic storage because of their relatively rapid dissolution rates.$^{13}$ In particular, the dissolution of divalent cation silicate minerals – such as those that contain Mg$^{2+}$ and Ca$^{2+}$ – is especially important because the leached cations can be used to “trap” anthropogenic CO$_2$ during a process known as
mineral trapping.\textsuperscript{4,7-10} For example, in underground mineral trapping, also known as \textit{in situ} storage, CO\textsubscript{2} is injected into a saline aquifer\textsuperscript{14} and reacts with water through reactions 1 and 2:

\begin{align*}
\text{CO}_2(\text{g}) + \text{H}_2\text{O}(\text{l}) & \rightarrow \text{HCO}_3^{-}(\text{aq}) + \text{H}^{+}(\text{aq}) & 1 \\
\text{CO}_2(\text{g}) + \text{H}_2\text{O}(\text{l}) & \rightarrow \text{CO}_3^{2-}(\text{aq}) + 2\text{H}^{+}(\text{aq}) & 2
\end{align*}

where the formerly gaseous CO\textsubscript{2} is now present as aqueous carbonate ions. The H\textsuperscript{+} formed during this transformation then reacts with the mineral surrounding the aquifer cavity. \textbf{Reaction 1-3} describes the case of a divalent cation silicate:

\[ \text{M}_2\text{SiO}_4(\text{aq}) + 4\text{H}^{+}(\text{aq}) \rightarrow 2\text{M}^{2+}(\text{aq}) + \text{Si(OH)}_4(\text{aq}) \]  

As the mineral dissolves, metal ions (M\textsuperscript{2+}) are released to solution and are available for reaction with the carbonate ions (CO\textsubscript{3}\textsuperscript{2-}), produced by the dissolution of CO\textsubscript{2} as described in \textbf{Reactions 1-1–2}. This interaction is described by the \textbf{Reaction 1-4}, showing the general precipitation scheme of carbonate mineral forms:

\[ \text{M}^{2+}(\text{aq}) + \text{CO}_3^{2-}(\text{aq}) \rightarrow \text{MCO}_3(\text{s}) \]

Thus, the gaseous CO\textsubscript{2} is “trapped” as a stable solid carbonate. Similar reactions can occur in \textit{ex situ} mineral trapping reactions where Mg-silicates are dissolved and the Mg\textsuperscript{2+} is used to precipitate Mg-carbonates.\textsuperscript{14} Despite the urgency for a means to store CO\textsubscript{2}, a molecular scale understanding of how these minerals release metal ions to solution does not exist; therefore, investigations into the breaking of M–O bonds as well as the distribution of protonation states of surface sites are required to provide insight into how metal ions are released from mineral surfaces.
1.2 Silicate Mineral Structures

In order to understand how metal ions are released from the mineral surface, one must begin by describing silicate mineral structures. The structure of each silicate mineral is based upon the SiO$_4^{4-}$ tetrahedron, where a Si atom is bonded to four O atoms (Figure 1-1a). Orthosilicate minerals are those where each SiO$_4^{4-}$ does not share any O atoms with the closest SiO$_4^{4-}$ within the mineral structure. The remaining mineral classes, however, share at least one apex of the tetrahedron with a neighboring SiO$_4^{4-}$ unit. Disilicate minerals share one mutual O atom between two SiO$_4^{4-}$ units. Single chain silicates, whose structure is shown in Figure 1-1b, share two mutual O atoms in each SiO$_4^{4-}$ tetrahedron. Double chain silicate minerals in Figure 1-1c share three O atoms per SiO$_4^{4-}$ tetrahedron, and this is akin to the single chain silicate structure in Figure 1-1b being joined together with its mirror. Ring silicate minerals have six SiO$_4^{4-}$ formula units linked together such that each tetrahedron shares two O atoms (Figure 1-1d). The structure for sheet silicate minerals is shown in Figure 1-1e, and here the ring silicate structure of Figure 1-1d is essentially extended laterally. Lastly, the framework silicate mineral structure is shown in Figure 1-1f, and in this case, every O atom within the SiO$_4^{4-}$ formula unit is shared between tetrahedra.
Figure 1-1: Formula units for silicate mineral structures. (a) Orthosilicate (b) Single chain (c) Double chain (d) Ring (e) Sheet (f) Framework.
In this work, the two main classes of silicate minerals discussed are orthosilicates (Figure 1-1a) and framework silicates (Figure 1-1f). The orthosilicate mineral that receives the most attention is the forsterite end-member of the olivine series. Forsterite, Mg$_2$SiO$_4$, has SiO$_4^{4-}$ tetrahedra that are connected via Mg$^{2+}$ ions. Each Mg$^{2+}$ is present in octahedral coordination such that the six O atoms from neighboring SiO$_4^{4-}$ tetrahedra satisfy this coordination. Two Mg$^{2+}$ ions for every SiO$_4^{4-}$ tetrahedron serve to balance the charge.\textsuperscript{16} These MgO$_6$ octahedra alternate the SiO$_4^{4-}$ tetrahedra in the mineral structure (Figure 1-2).\textsuperscript{17}

In addition to forsterite, anorthite (CaAl$_2$Si$_2$O$_8$) is discussed throughout this work. The anorthite end-member of the plagioclase group of framework silicates is a Ca-aluminosilicate. In this mineral, both Si and Al are tetra-coordinated to four O atoms, forming SiO$_4$ and AlO$_4$ tetrahedra (Figure 1-3).\textsuperscript{18} Four tetrahedra are bound where each shares two O atoms with its neighbor, forming a ring. Rings are bound together by sharing additional O atoms and are arranged in spirals. The “framework” arises from the lateral joining of these rings. The resulting structure is open and therefore can accommodate larger cations such as Ca$^{2+}$.\textsuperscript{12} Structural analysis of the Ca$^{2+}$ within the mineral shows that it is likely hepta-coordinated.\textsuperscript{19}
Figure 1-2: Forsterite mineral structure from the Materials Studio mineral structures database.\textsuperscript{18} The Mg\textsuperscript{2+} ions are yellow, the Si tetrahedra are green, and the O atoms are red.

Figure 1-3: Anorthite mineral structure.\textsuperscript{18} The O atoms are red, the Si tetrahedra are green, the Al\textsuperscript{3+} tetrahedra are blue, and the Ca\textsuperscript{2+} ions are cyan.
1.3 Kinetics

1.3.1 Transition State Theory (TST)

The transition state theory (TST) approximation is used to describe the rate of dissolution for the divalent cation silicates discussed herein. TST describes reaction mechanisms and reaction rates and hinges upon the existence of a short-lived chemical species that has characteristics of both the reactants and the products of the reaction. TST is derived from the statistical nature of the reacting species but, in its most classical state, does not include quantum contributions. The change from reactants to products is traced along the reaction coordinate, and the transition state corresponds to the structure with the highest free energy along that path. The transition state (TS) is more specifically a saddle point where the maximum energy along the reaction coordinate serves as the minimum energy point along the path orthogonal to the reaction coordinate (Figure 1-4).20

There are several assumptions made by the TST that become critical when calculating the rate of a reaction. First, the TST incorporates the Born-Oppenheimer approximation, where electronic motion is decoupled from the nuclear motion. Second, a Maxwell-Boltzmann distribution exists among reactant molecules. Third, those reactants that contain sufficient energy to cross the TS barrier toward the products are not able to recross this barrier and return to reactants (i.e. the reaction is irreversible). Fourth, movement along the reaction coordinate is independent from other molecular motions. Lastly, those TSs moving toward products exist in states according to Maxwell-Boltzmann distributions.20
The equilibrium that is assumed to exist between the reactants and the TS is known as the quasi-equilibrium hypothesis. In this description, a model reaction is considered according to **Reaction 1-5**:\[ A + B \rightarrow X^\dagger \rightarrow C \] where A and B are the reactants, $X^\dagger$ is the transition state, and C is the product. The energy profile of this model reaction is shown in Figure 1-4. Species moving toward the products proceed from left to right in the plot in Figure 1-4, and those moving from products to reactants move from right to left. The TS exists at the center of the plot where the potential energy is the greatest. This region is known as the dividing surface and is used to delineate those species that have “crossed over” from reactants to products or vice versa. Thus TSs may either be those species forming as products become reactants or as reactants become products. The rate of the reaction describes the speed at which TSs are converted to products (forward) or reactants (reverse).\[ ^{20} \]
1.3.2 Rate Laws

From the classical TST, the rate for a reaction can be calculated using the energy difference between the reactants and TS. The difference in internal energy between the reactants and the TS $\Delta U_{\ddagger}^{\circ}$ can be related to the activation energy $E_a$ according to Equation 1-6:

$$E_a = RT + \Delta U_{\ddagger}^{\circ}$$

where $R$ is the gas constant and $T$ is the temperature in K. $E_a$ is defined as the minimum energy required for a reaction to occur and can be used to calculate the rate constant $k$ for a reaction through the Arrhenius equation (Equation 1-7):

$$k = A e^{-E_a/RT}$$

where $A$ is the Arrhenius pre-exponential factor. For the gas phase, $A$ is defined by Equation 1-8:

$$A = e^{-(\Delta n\downarrow - 1)} \frac{k_B T}{h} e^{\Delta S_{\ddagger}^{\circ}/R}$$

where $\Delta n\downarrow$ is the change in moles between the reactants and TS, $k_B$ is Boltzmann’s constant, $h$ is Planck’s constant, and $\Delta S_{\ddagger}^{\circ}$ is the standard entropy change between the reactants and TS.

Several factors affect the rate constant; however, $E_a$ and $T$ are of particular interest to this work. Due to the inverse exponential relationship between $k$ and $E_a$, a greater $E_a$ is associated with a smaller $k$. The temperature of a reaction, however, has the opposite effect on $k$ such that both increase concurrently as described in Equation 1-7.
In addition, the temperature dependence of \( A \) contributes to the temperature dependence of \( k \).\(^2^0\)

Reaction rates are not only affected by temperature but on the relative concentrations of reactants. For example, a zero-order reaction has no concentration dependence,\(^2^0\) and the relationship between the rate of the reaction and the concentration of a reactant follows **Equations 1-9–10:**

\[
r = k[A]_0 = k 
\]

where \( r \) is the rate of the reaction, \( k \) is the rate constant, and \( [A]_0 \) is the initial concentration of the reactant.\(^2^2\) **Equation 1-9** can be integrated to yield the slope-intercept form of a line:

\[
[A]_t = -kt + [A]_0 
\]

where \( [A]_t \) is the concentration at time \( t \). **Figure 1-5** shows a typical zero-order reaction, where the plot follows **Equation 1-10**, and the rate constant \( k \) is the slope of the line.

However, if the rate of the reaction increases linearly with the concentration of a reactant, then the reaction is of the first order and is described by **Equation 1-11**:

\[
r = k[A] 
\]

**Equation 1-11** can be integrated and rearranged to give slope-intercept form, shown by **Equation 1-12**:

\[
\ln[A]_t = -kt + \ln[A]_0 
\]

A plot of **Equation 1-12** is given in **Figure 1-6**.
In some cases, the rate of a reaction increases with the square of the concentration of a reactant, and these reactions are known as second-order. The rate law for a second-order reaction is defined as Equation 1-13,

\[ r = k[A]^2 \]  

and is integrated to form Equation 1-14:

\[ \frac{1}{[A]_t} = kt + \frac{1}{[A]_0} \]  

Figure 1-7 displays a second-order reaction. 

The orders of reactions can be classified according to the effect of each individual reactant on the overall rate of a reaction. If, for example, a rate doubles when [A] doubles, then the reaction is first-order with respect to A. However, if the rate of the same reaction quadruples as [B] doubles, then the reaction is second-order in B. The overall order of a
reaction is the sum of the orders of the individual reactants, and in this example, the reaction is third-order overall.\textsuperscript{20}

For reactions involving more than one step\textsuperscript{20} or where one reactant is present in great excess of another,\textsuperscript{22} the pseudo-first order approximation can be used. In
experimental situations where \([A] >> [B]\),\(^{20}\) the rate of the reaction appears to behave independently of \([A]\).\(^{22}\) For those reactions where bulk water serves as a reactant, the pseudo-first order approximation is invoked.\(^{22}\) Some geochemical reactions are examples of those that appear as pseudo-first order reactions and will discussed in Section 1.4.

Computational determinations of experimental rate laws have recently been employed to describe geochemical systems.\(^{23,24}\) In these investigations, molecular clusters representing single surface sites were used, and the energy required to break the Si–O bond\(^{23}\) bond and Mg–O bond\(^{24}\) was calculated. This energy was used to calculate the rate constant as in Equation 1-7, and the rate laws for dissolution of quartz\(^{23}\) and forsterite\(^{24}\) were described according to the pseudo-first order approximation.

### 1.4 Geochemical Reactions

Geochemical reactions are studied for their contributions to natural\(^{11}\) and energy-related processes,\(^{4-10}\) but understanding the kinetics of these reactions is challenging for a number of reasons. First, these reactions often occur in aqueous media, and therefore, water as well as dissolved species are reacting with the mineral under investigation, leading to complex competing or side reactions. Further, separating the contributions of water, \(\text{H}_3\text{O}^+\), \(\text{OH}^-\), and additional ions such as \(\text{Na}^+\), \(\text{Cl}^-\), or \(\text{Ca}^{2+}\) compounds this issue, as dissolved ions are known to catalyze some geochemical reactions.\(^{25-29}\) In addition, determining the concentrations of reactants may often be difficult.\(^{30-33}\) For example, quantifying the number of sites on a surface that participate in a given reaction remains poorly understood since many investigations have focused on the total number of surface
sites as opposed to only those sites that participate in the reaction.\textsuperscript{32-34} For dissolution, the release of species to solution is considered to be pseudo-first order because of the excess of water in the system, but the rate law for each mineral differs\textsuperscript{34} and depends strongly upon pH.\textsuperscript{35-38} Each effect on dissolution must be investigated individually to determine its impact on the rate, and there are a number of experimental and theoretical approaches that can be used, which will be discussed below.

1.4.1 Dissolution Reactions

Dissolution of a mineral can be defined as the collection of physicochemical phenomena that cause the ultimate breakdown of the mineral structure and lead to the net release of elemental and ionic constituents to solution.\textsuperscript{39} If one traces these phenomena chronologically, then a complete picture of dissolution can be described. First, species from solution are transported to the mineral surface, and these include H\textsubscript{2}O molecules, H\textsubscript{3}O\textsuperscript{+} ions, OH\textsuperscript{-} ions, and dissolved species such as organic acids, metal cations, and halide anions.\textsuperscript{39} Next, these species adsorb to the mineral surface, a process thought to contribute to degradation of the mineral surface.\textsuperscript{40} The adsorption of ions,\textsuperscript{39,41,42} such as H\textsubscript{3}O\textsuperscript{+}, to the surface O atoms weakens the M–O bond, allowing this bond to break more easily.\textsuperscript{39-41} Additional chemical reactions lead to the alteration of the mineral surface structure, such as the development of surface defects. These features are sites where surface reactions can preferentially occur.\textsuperscript{43,44} As mineral constituents are released to solution, additional ions from solution replace them in the mineral structure to allow for charge-balancing.\textsuperscript{45} The final step of dissolution is the transport of dissolved species.
away from the aqueous-mineral interface into solution. Figure 1-8 shows the dissolution rate of anorthite as pH increases, and such an analysis is typical of experimental descriptions of dissolution. Both experimental and theoretical approaches to investigating dissolution focus on examining each of these processes in turn, and these techniques and models will be discussed in the following section.

1.4.2 Methods for Studying Dissolution

Dissolution of geochemical materials can either be studied experimentally or computationally. The rates of these reactions are determined by measuring the concentration of product species in solution over time, and thus, these experiments rely on the release of chemical species from the mineral surface to solution. Once the rate of a dissolution reaction has been determined, the rate constant for a reaction, and subsequently the activation energy, may be determined. Computational methods approach dissolution reactions oppositely such that the energy for a given reaction is calculated, and the rate constant is calculated using Equation 1-7. The goal of a computational approach investigating dissolution is the ability to either compare observed trends in experimental data or to calculate a quantity that is experimentally measured. In the two sections below, various types of experimental and computational approaches to studying dissolution reactions are described.

In the rate law for a geochemical reaction, the surface area of the mineral sample is experimentally measured and is assumed to be proportional to the reaction rate.\textsuperscript{34-38}
One of the most common methods for determining the surface area of a mineral sample is the Brunauer-Emmett-Teller (BET) surface area.\textsuperscript{47} In this approach, an inert gas adsorbs to the mineral surface, and the surface area of the sample is determined according to the volume of gas that was lost to adsorption. The surface area that is calculated is then inserted into the rate law and used to explain the contribution of surface sites to the reaction rate. Unfortunately, the BET surface area does not always correlate to the dissolution rate of mineral samples,\textsuperscript{32} resulting in the necessity to quantify the reactive surface sites needed to describe the surface contribution to dissolution.\textsuperscript{32,48} That is to say, instead of considering the entire surface area as participating in dissolution, only individual sites that are known to be more reactive than others are considered to play a role in dissolution.\textsuperscript{32,48}

Computational descriptions of mineral surface samples can provide insight into the problem of total surface sites versus reactive surface sites. Modeling the surface topography shows where adsorption of gas particles to the surface is most likely and also

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**Figure 1-8:** Anorthite dissolution versus pH. The various shapes represent the input concentrations of Si, Ca\textsuperscript{2+}, and Al\textsuperscript{3+}.\textsuperscript{46}
provides insight into the role of surface structure on the adsorption of gas particles.\textsuperscript{44,49} Because dissolution is comprised of a collection of processes,\textsuperscript{39} computational approaches using molecular clusters focus on representing individual sites to explain the contribution of each to reactions of dissolution on the mineral surface,\textsuperscript{23,24,26,27,50-54} and these calculations further provide insight into which steps are rate-limiting.\textsuperscript{24}

1.4.2.1 Experimental Methods

Several experimental approaches have been used to study the dissolution of minerals. The two that have the most popularity are batch reactors\textsuperscript{5,10,33,35,45,55} and flow-through reactors.\textsuperscript{8,32,34,40,43,56} Batch reactors consist of a single container where all of the reactants are placed together and continuously stirred to promote homogeneous reaction conditions in solution. The reaction progress is measured by the accumulation of products in solution, and the concentration can be followed with time. The advantage of this method is that these reactors are easily constructed. Unfortunately, frequent sampling of the solution leads to a complicated description of the reaction rate, particularly if the rate changes with time.\textsuperscript{57} Batch reactors have been used to study the dissolution of a variety of minerals, such as alumina,\textsuperscript{41} basalt,\textsuperscript{36} beryllium oxide,\textsuperscript{41} forsterite,\textsuperscript{5,35,45,55} kaolinite,\textsuperscript{33,38} montmorillonite,\textsuperscript{33} serpentine,\textsuperscript{10} and silica.\textsuperscript{28}

The other commonly used experimental method for dissolution is a flow-through reactor, also known as a mixed-flow reactor. In this setup, fluid moves through the reaction vessel at a continuous rate. The concentration of products in the outgoing solution is monitored and used to gauge the progress of the reaction. The advantage to
this method is that the reaction rate is determined directly as the concentration of products increases with time, but the disadvantage is that these apparati require difficult construction.\textsuperscript{57} Mixed-flow reactors have been used to study the dissolution of aluminosilicates,\textsuperscript{32} anorthite,\textsuperscript{34} forsterite,\textsuperscript{8,40,56} and quartz.\textsuperscript{25,43}

Two other less commonly used experimental approaches for studying dissolution are surface titrations\textsuperscript{30,31} and the pH stat method.\textsuperscript{42,58,59} In a surface titration experiment, the mineral sample is suspended in solution, and either acid or base is added step-wise.\textsuperscript{30} The pH is monitored throughout the experiment, and the loss of either $\text{H}_2\text{O}^+$ or $\text{OH}^-$ ions from solution indicates adsorption to the mineral surface.\textsuperscript{30,31} The pH stat method is similar to a batch reactor where the reaction is contained in a single vessel, and the solution is sampled periodically.\textsuperscript{59} A constant pH is maintained by the addition of acid to compensate for $\text{H}^+$ ions used in the dissolution reaction scheme.\textsuperscript{59}

The strength of experimental approaches, then, is the ability to follow dissolution processes simultaneously and measure the concentration of several products or the effect of several environmental factors within the same reaction scheme. Despite the existence of several experimental methods for studying the dissolution of minerals, developing an accurate molecular-scale picture of the processes occurring in solution at the mineral surface is still an elusive challenge in these studies. In order to understand geochemical processes from a molecular perspective, computational methods have become another tool used to probe the nature of these systems.
1.4.2.2 Computational Methods

The strength of computational methods is the ability to describe a molecular-scale picture of an individual process within a geochemical reaction scheme. Several temporal and spatial scales are possible using computational tools, and these regimes are depicted in Figure 1-9. The challenge, however, is to include enough chemical information so the calculations provide molecular information while simultaneously representing a system size or timescale that provides meaningful insight. Further, this challenge is compounded by the size of the system that can be studied. Smaller systems are required to include chemical information such as electronic structure, and these calculations are tractable up to ~50 atoms. Larger systems of $10^2 - 10^5$ atoms can also be modeled, but the tradeoff is that the electronic structure is not included, which permits these calculations to remain feasible.

The smallest-sized systems can be studied with electronic structure calculations, and one particular class of electronic structure calculations employs the density functional theory (DFT). In this approach, the electronic structure of atoms within the system is modeled with approximations to the Schrödinger equation, and these approximations are manifested via functionals that use the energy and electron density gradients to model electronic behavior. These functionals include several quantities to describe the energy and electron density. The exchange-correlation functional combines the energy of the inversion of two electrons within the wavefunction with the energy arising from the coupled motion of electrons throughout the system. The exchange-correlation functional can be further modified by addition of Hartree-Fock
exchange, where electron exchange is described according to Hartree-Fock theory.\textsuperscript{64} An
advantage of DFT is that each electron has a maximum of four coordinates: x, y, z, and spin direction, while the electron density is described solely in terms of three coordinates, regardless of how many electrons are present within the system. The use of density functional theory (DFT) calculations has recently been reviewed,\textsuperscript{65} and the functional used throughout this work blends exchange-correlation with Hartree-Fock exchange.\textsuperscript{64-68}

DFT calculations have been employed to model geochemical reactions.\textsuperscript{23,24,26,27,50-54}

In order for these electronic structure calculations to be practical, model clusters are designed to effectively represent the system under study. Once a proper cluster has been chosen, these structures are energy-minimized to isolate a stable molecular geometry.
Clusters to represent the reactants, transition state, and products of a reaction are compared to describe the reaction mechanism, and the relative energies of the reactants and transition state give the barrier height for the reaction. The descriptions of the electronic structure within these calculations are derived from the gas-phase. However, the aqueous-phase can be modeled through the inclusion of a dielectric continuum or a super-molecule, where explicit H₂O molecules around the reacting species are included.

Increasing the system size from molecular clusters to a unit-cell of a crystal can be attained via density functional theory molecular dynamics (DFT-MD) simulations. The electronic structure of atoms within the simulation cell is included, and motion of the atoms is also permitted. The electronic structure is represented by pseudo-potentials that divide the electrons into core and valence (i.e. non-reacting and reacting), and the band structure is evaluated via a projector-augmented wave (PAW) approximation.

The PAW approximation is an all-electron approach that includes full wavefunctions and blends augmented waves with pseudopotentials. The augmented wave methods serve to fraction the electronic system into atom regions and envelope functions, which describe bonding between atoms. The electrons in the atom region are fractioned into valence and core, and the projector functions serve to describe the local nature of wavefunctions within the atomic region. Despite these adjustments to evaluating electronic structure, chemical behaviors using all-electron methods are well replicated.

DFT-MD simulations provide an opportunity to investigate reactive processes and bulk characteristics within one simulation scheme, and one application for which this tool is particularly suited is the study of the aqueous-mineral interface. Reactions between
atoms are possible, and the inclusion of explicit H$_2$O molecules in solution permits representation of bulk water. Previous applications of these simulations for geochemical systems include modeling the adsorption of H$_2$O molecules to mineral surfaces,\textsuperscript{75} the aqueous-mineral interface,\textsuperscript{76,77} the adsorption of ions from solution to the mineral surface,\textsuperscript{78} and the vibrational spectrum of H$_2$O molecules adsorbed to the mineral surface.\textsuperscript{79}

Monte Carlo (MC) simulations are practical for system sizes of hundreds\textsuperscript{80-86} to thousands of particles.\textsuperscript{87-90} The fundamental design of this approach relies on the relative energies of the arrangement of particles in the system as they are moved in accordance with the simulation scheme under study. When the present arrangement is lower in energy than the initial, the system accepts the present arrangement. If instead the initial arrangement is lower, then two options are possible. The Boltzmann factor, $e^{-\Delta E/k_BT}$, is compared with a number between 0 and 1. If the Boltzmann factor is less than this fraction, the present arrangement is assumed. If on the other hand, the Boltzmann factor is greater than this fraction, then the initial arrangement of particles in the system remains.\textsuperscript{60} This approach allows for the equivalent sampling of all possible configurations of the system, where the probability is weighted by the Boltzmann factor.\textsuperscript{91} Because MC simulations do not include a time parameter,\textsuperscript{92} those phenomena that proceed over a long timescale (i.e. ns or longer) can be studied. Geochemical processes such as the H-bonding networking on the mineral surface,\textsuperscript{86} adsorption of water to the mineral surface\textsuperscript{80,84,85,90} or to the interlayer,\textsuperscript{81-83} evolution of surface topography,\textsuperscript{88}
abiotic dissolution and precipitation, biologically-mediated dissolution, and pore-water fluid flow have each been studied via MC simulations.

The last class of computational methods is molecular dynamics (MD) simulations, and this approach is feasible for systems similar in size to those for MC simulations. MD simulations use Newton’s second law to describe the interaction of particles, and their behavior is studied with time. The electrostatic interactions of particles are modeled with potentials, which is in contrast with electronic structure and DFT-MD methods where the electronic structure of each particle is explicitly included. Specific characteristics of electronic behavior such as vibrational motion, Coulombic and Lennard-Jones interactions, and van der Waals interactions are also present in these simulations. Recent applications of MD simulations to geochemical systems include examining the silica-water interface as a function of surface hydroxylation, the forsterite structure under high temperature and pressure conditions, the adsorption of water to the forsterite surface, and pore water fluid flow.

The two computational methods that are employed in this work are density functional theory (DFT) calculations and DFT-MD simulations. DFT calculations are used to investigate the breaking of M–O (M = Mg\(^{2+}\), Ni\(^{2+}\), and Ca\(^{2+}\)) bonds within silicate molecular clusters to model the release of these metal ions from mineral surfaces. DFT-MD simulations are used to model the aqueous-mineral interface for forsterite (100) and (010) surfaces, and reactions between surface groups and between surface groups and species from solution permit a realistic model of the forsterite surface structure in an aqueous environment.
1.5 Expected Implications

The expected implications of this work are to provide a molecular scale understanding of the mechanisms through which dissolution reactions occur and to investigate contributing factors to experimental values. In particular, the barrier heights of metal release from mineral surfaces are calculated, and experimental data are combined with computational results to calculate rates and rate constants of dissolution. These two techniques relate to the need to understand how Mg$^{2+}$ and Ca$^{2+}$ are released from forsterite and anorthite during CO$_2$ storage via mineral trapping. The functional groups present on the forsterite surface at the water-mineral interface are also modeled, and this investigation provides insight into which sites on the surface are available to participate in dissolution reactions. These approaches are taken to explain experimental observations such as the dissolution rates of forsterite in Chapter 2, the dissolution rates and water-exchange rates of Ni-, Mg-, and Ca-silicate minerals in Chapter 3, and the difference between the stabilities and functional groups present on the forsterite (100) and (010) surfaces in Chapter 4. The use of computational techniques to study geochemical systems employs existing technology and relies upon models to adequately explain experimental phenomena. The work described is particularly unique for its inclusion of alkaline and transition metals as well as the methods chosen to model experimental observations.
1.6 References


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Chapter 2

A Description of Mg$^{2+}$ Release from Forsterite using Ab-Initio Methods

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2.1 Introduction

The work described in Chapter 2 employs quantum mechanical calculations to model the release of Mg$^{2+}$ from sites on the forsterite mineral surface. Molecular cluster analogs of these surface sites allow for tractable computation times, and the varying protonation states of sites are included. The reaction mechanism and the barrier height for Mg$^{2+}$ release from each type of site are reported. The barrier heights are used to calculate rate constants for Mg$^{2+}$ release from each type of site, and these are combined with experimental data to calculate a rate for Mg$^{2+}$ release from the forsterite surface versus pH.

In order to understand the dissolution of forsterite, one begins with a description of the forsterite mineral structure (Figure 2-1). The bulk structure of forsterite has octahedrally coordinated Mg bound to six oxygen atoms, each of which is bound to a Si atom. The network continues, alternating between tetrahedrally-coordinated Si and octahedrally-coordinated Mg atoms connected via bridging oxygen atoms (O).$^{19,20}$ The release of Mg species to solution is triggered by the breaking of a terminal Mg–O bond linking the Mg to the surface, and in this state, Mg is octahedrally-coordinated with the O
Figure 2-1: Forsterite (100) surface where a Mg–O–Si surface site has been extracted to depict the molecular cluster model. The magnesium atoms are yellow, the oxygen atoms are red, the silicon atoms are turquoise, and the hydrogen atoms are white.

and five H₂O molecules as is shown by the molecular cluster in the inset to Figure 2-1. Once released from the surface, Mg returns to six-fold coordination when an incoming H₂O molecule replaces the O for the sixth position around the Mg center. Descriptions of this process based on empirical studies are both numerous and extensive and are outlined below.

In a recent review,²¹ the dissolution rates of a number of silicate minerals, including forsterite, were compared as a function of pH. For silica, the dissolution rate is at a minimum at neutral pH and increases in acidic and basic pH ranges, but for forsterite, most recent work⁵,⁷,¹¹,²¹ supports a linear decrease in the Mg²⁺ release rate with an increase in pH as shown in Figure 2-2.¹² The experiments performed by Wogelius and Walther⁶ and Blum and Lasaga²² showed an increase of the dissolution rate in the basic
region. Blum and Lasaga explained that their U-shaped dissolution rate curve analogous to that observed for silica may indicate that the deprotonation of Si sites, defined here as Si–O–Mg sites, on the forsterite surface is the most prevalent hydrolysis surface reaction at basic pH. However, the validity of the studies by Wogelius and Walther\textsuperscript{6} and Blum and Lasaga\textsuperscript{22} have been questioned because neither set of experiments was performed in the steady-state region of dissolution nor was a statistically adequate number of data collected for the dissolution rate in the basic pH region.\textsuperscript{8} Further, the ionic strength of the solutions that Wogelius and Walther used during their dissolution studies of forsterite could have produced an increase in surface charge\textsuperscript{23} which may lead to an increase in dissolution rate.\textsuperscript{24,25}
A recent study by Rosso and Rimstidt\textsuperscript{9} investigated the effect of H\textsubscript{3}O\textsuperscript{+} adsorption onto the forsterite surface in the pH range of 1 to 3, and the link between H\textsubscript{3}O\textsuperscript{+} catalysis and forsterite dissolution was explained. They determined that two H atoms from H\textsubscript{3}O\textsuperscript{+} adsorb to O atoms in two adjacent Mg octahedra which weakens the Mg–O bonds. This description explains the observation that two H\textsuperscript{+} replace Mg\textsuperscript{2+} in the surface structure during dissolution.\textsuperscript{20} Further, this allows H\textsubscript{2}O molecules from solution to more easily exchange for the O atoms in the mineral structure, and the substitution causes the surface network to open. After five H\textsubscript{2}O molecules have replaced the O atoms in the octahedra, the Mg is hexa-coordinated to five H\textsubscript{2}O molecules and one O, and this last step before release to solution\textsuperscript{26} is amenable to a molecular-scale interrogation of the forsterite dissolution process.

In order to explain experimental observations with a molecular-level understanding, ab-initio calculations have recently been employed to describe mechanisms for hydrolysis reactions occurring at the mineral surface.\textsuperscript{2,4,24,25} In these studies, molecular cluster-sized analogs of mineral units are used to represent reaction sites on the surface, and this approach has replicated experimental trends for both silica\textsuperscript{2} and aluminosilicate minerals.\textsuperscript{4} The use of molecular clusters enables the description of the central phenomenon in a chemical process without the computational burden of atoms that do not contribute directly to the reaction in question.\textsuperscript{24} This facilitates a molecular description of dissolution where the molecular cluster serves as a site on the surface, and H\textsubscript{3}O\textsuperscript{+}, H\textsubscript{2}O, or OH\textsuperscript{−} is the second reactant. At every pH, a distribution of each type of site is present.
One of the challenges in choosing an ab-initio approach to study geochemical reactions is that the calculations are often performed in the gas-phase and thus with no solvent explicitly present, while the reactions occur at the water-surface interface in aqueous media. Including a solvation continuum model is arguably necessary because it would yield insight into how effectively the gas-phase calculations represent the chemical process under study, and the inclusion of solvation effects has recently been employed in the investigation of species with geochemical relevance.\textsuperscript{27,28} In addition, comparisons of ab-initio results with experimental data are more meaningful when solvation effects are included because they allow for the identification of any results which are artifacts of performing the original calculations in the gas-phase. The use of a continuum model is one tool for the inclusion of a solvation approach without the use of explicit H\textsubscript{2}O molecules.\textsuperscript{27,29}

The aim of this study is to examine dissolution from Mg\textsuperscript{2+} sites, defined as Mg–O–Si sites, on the forsterite surface using a molecular cluster approach via ab-initio calculations. The reaction mechanisms are described along with energy profiles that present the barrier heights for these reactions in gas-, single-point aqueous-, and energy-minimized aqueous-phases. Rate constants for these reactions as well as the rate for the release of Mg\textsuperscript{2+} from Mg–O–Si sites are also presented. These computational results are compared with experimental data.

### 2.2 Computational Methods

Molecular clusters were extracted from Mg–O–Si sites on the forsterite (100) surface using the Cerius\textsuperscript{2} database,\textsuperscript{30} and an example of a cluster is shown in the inset of
**Figure 2-1.** Here a Mg$^{2+}$ is bonded to a O which is in turn bonded to a Si atom. The Si atom is bonded to three OH groups which are meant to represent the bonds to the bulk crystal, and this is in accord with the finding that bonds breaking at the surface experience minimal contribution from other atoms deeper in the surface.\textsuperscript{24} The hexacoordination of the Mg$^{2+}$ is satisfied by five H$_2$O molecules in its first hydration sphere, and the cluster is reacted with a H$_2$O molecule to simulate dissolution. **Figure 2-3** depicts a model molecular cluster with a H$_2$O molecule which represents a general setup for each of the calculations presented here. In each of these clusters, the protonation or deprotonation of a surface site is also considered to represent acidic and basic conditions, respectively. The presence of an H$^+$ on the O represents a protonated site, and for the neutral and deprotonated sites, no H$^+$ is included. In the deprotonated site reaction, the absence of an H$^+$ on the surface site is represented by reacting the molecular cluster with an OH$^-$ rather than a H$_2$O molecule.

The hydrolysis of Mg–O–Si sites in each protonation state is represented by the following reactions, corresponding to protonated, neutral, and deprotonated Mg–O–Si sites, respectively. The protonated Mg–O–Si site reaction is (2-1):

\[
[Mg(H_2O)_5(OH)Si(OH)_3]^{2+} + H_2O \rightarrow [Mg(H_2O)_6]^{2+} + Si(OH)_4
\]  

where presence of an H$^+$ on the O denotes a protonated site. The neutral Mg–O–Si site reaction is described by two possible reactions (2-2):

\[
[Mg(H_2O)_5OSi(OH)_3]^+ + H_2O \rightarrow [Mg(H_2O)_5(OH)]^+ + Si(OH)_4
\]  

and (2-3):

\[
[Mg(H_2O)_3OSi(OH)_3]^+ + H_2O \rightarrow [Mg(H_2O)_3(OH)]^+ + Si(OH)_4
\]
Figure 2-3: The model molecular cluster used to represent Mg–O–Si sites on the forsterite surface. The color scheme is the same as Figure 2-1.

\[ [\text{Mg(H}_2\text{O)}_5\text{OSi(OH)}_3]^+ + \text{H}_2\text{O} \rightarrow [\text{Mg(H}_2\text{O)}_6]^2^+ + \text{SiO(OH)}_3^- \]

where the O is not protonated, and the products may include either a hydroxylated surface site, denoted by Si(OH)$_4$, or a deprotonated surface site, denoted by SiO(OH)$_3^-$. The deprotonated Mg–O–Si site reaction is also described by two possible reactions (2-4):

\[ [\text{Mg(H}_2\text{O)}_5\text{OSi(OH)}_3]^+ + \text{OH}^- \rightarrow [\text{Mg(H}_2\text{O)}_3\text{(OH)}_2] + \text{H}_2\text{O} + \text{Si(OH)}_4 \]

and (2-5):

\[ [\text{Mg(H}_2\text{O)}_5\text{OSi(OH)}_3]^+ + \text{OH}^- \rightarrow [\text{Mg(H}_2\text{O)}_4\text{(OH)}]^+ + \text{H}_2\text{O} + \text{SiO(OH)}_3^- \]

where the O is not protonated as with the neutral site reaction, and the products may include either a hydroxylated surface site, denoted by Si(OH)$_4$, or a deprotonated surface site, denoted by SiO(OH)$_3^-$. In each calculation, the overall charge of the system is included, and the contribution of each species to the overall charge is shown in Reactions 2-1, 2-2, 2-3, 2-4, and 2-5.
A density functional theory (DFT) approach was chosen for this study because of enhanced computational feasibility as well as successful use of these methods in the literature to calculate the hydrolysis reaction mechanisms for mineral dissolution.\textsuperscript{2,4,24,25} The B3LYP functional is a hybrid density functional which includes exchange-correlation,\textsuperscript{31-34} gradient correction,\textsuperscript{33,34} and a fraction of Hartree-Fock exchange.\textsuperscript{35} The employment of the 6-31+G(d,p) basis set is reasonable in this application because it is appropriate for Mg\textsuperscript{2+} complexes,\textsuperscript{36} and \textit{p} orbitals were added in the event that a H\textsuperscript{+} transfer occurred in the course of any of the reactions studied.

The structures of the reactants for each reaction were optimized without constraints in the gas-phase, and then potential energy surface (PES) scans of the Mg–O distance were performed to determine estimates for both the transition states and products which were in turn optimized. Reactant and product species had no negative frequencies, and possible transition states were identified by the presence of one negative frequency that corresponded to the breaking of the Mg–O bond. A second set of PES scans was performed where the distance between the incoming H\textsubscript{2}O molecules and the Mg\textsuperscript{2+} was decreased in order to calculate the barrier height for the absorption of H\textsubscript{2}O by Mg\textsuperscript{2+}. The energy profiles were constructed by using the reactant energy as the reference state and comparing the energies of the transition states and products to the energy of the reactants. All calculations were performed with Gaussian 03,\textsuperscript{37} and all molecular cluster figures were visualized with GaussView 4.\textsuperscript{38}

After the initial reaction mechanism was determined using gas-phase calculations, aqueous-phase calculations were performed using the self-consistent reaction field (SCRF) method through the integral equation formalism polarized continuum model.
with water as the solvent. Single-point aqueous-phase calculations were performed where the gas-phase structures were used as input, and the corresponding energies were calculated. No frequency calculations were performed because the gas-phase structures do not correspond to a potential energy minimum once the continuum solvation model is applied.

Finally, the reactants from the gas-phase calculations were optimized without constraints using the IEFPCM, and PES scans of the Mg–O distance as well as the approach of the H₂O to the Mg²⁺ were performed in a manner analogous to the gas-phase calculations. The reactants and products for each reaction yielded no negative frequencies, whereas optimized transition states in the aqueous-phase calculations had a single negative frequency corresponding to the breaking of the Mg–O bond.

In an effort to determine whether each hydrolysis reaction is spontaneous, the Gibbs free energy change for each reaction was calculated. In the gas-phase, the Gibbs free energy of each structure includes a zero-point energy (ZPE) correction as well as a correction to the thermal energy. Partition functions that describe the translational, electronic, rotational, and vibrational motion are incorporated into the entropy term, while the enthalpy is the atomic energy of the structure with the addition of a thermal correction factor.

In the continuum calculations, the Gibbs free energy is calculated with respect to contributions from the presence of the solvent continuum and the solute structure in the gas phase. The Gibbs free energy includes this electrostatic description of the solute/solvent interaction and a “cavitation free energy” contribution, which calculates the amount of energy required to form the cavity and includes the surface tension, the
surface area of the solute cavity, and the ratio of the molar volume of the solute to molar volume of the solvent. The temperature is treated as 298.15 K, and the free energy from the reaction was determined as (2-6):

$$\Delta G_{\text{reaction}} = \Delta G_{\text{products}} - \Delta G_{\text{reactants}}$$

where $\Delta G_{\text{reaction}}$ is the Gibbs free energy for the reaction, $\Delta G_{\text{products}}$ is the Gibbs free energy for the products, and $\Delta G_{\text{reactants}}$ is the Gibbs free energy for the reactants.

The Gibbs free energy quantities from the aqueous-phase calculations on the gas-phase structures relied upon additional corrections, as some of the correction quantities are calculated for the gas-phase only. Thus, the Gibbs free energy for the single-point aqueous-phase calculations was calculated according to (2-7):

$$\Delta G_{\text{reaction}} = (\Delta G_{\text{products}} + C_{P_{\text{gas-phase}}}) - (\Delta G_{\text{reactants}} + C_{R_{\text{gas-phase}}})$$

where $C_{P_{\text{gas-phase}}}$ is the Gibbs free energy correction from the gas-phase for the products and $C_{R_{\text{gas-phase}}}$ is the Gibbs free energy correction for the reactants, and this approach was used because the gas-phase structures were used in these calculations.

The Gibbs free energy for the reactions using optimized aqueous-phase structures was determined as (2-8):

$$\Delta G_{\text{reaction}} = (\Delta G_{\text{products}} + C_{P_{\text{aqueous-phase}}}) - (\Delta G_{\text{reactants}} + C_{R_{\text{aqueous-phase}}})$$

where $C_{P_{\text{aqueous-phase}}}$ is the correction to the products in the aqueous-phase and $C_{R_{\text{aqueous-phase}}}$ is the correction to the free energy of the reactants in the aqueous-phase. The Gibbs free energy values for the reactions in gas-phase, single-point aqueous-phase, and optimized aqueous-phase were used to determine if each reaction is spontaneous.
Once the reactant and transition state had been optimized in the gas- and aqueous-phases for each reaction, the frequency output files were used to calculate the rate constant using TheRate via Equation 2-9:

$$k_T = AT^n e^{-\frac{E_a}{RT}}$$

where $A$ is the pre-exponential factor, $T$ is the temperature in K, $n$ is the exponential order of the temperature-dependence of $A$, $E_a$ is the activation energy, and $R$ is the gas constant. Transition state theory (TST) was employed, and the rate constant was calculated at 298K. The value of $n$ was set to 0.0010 to eliminate the temperature dependence of $A$. For the single-point aqueous-phase calculations, the barrier heights were inserted into Equation 2-9 using the $A$ and $n$ quantities from the calculations for the gas-phase reaction because the structures were the same.

The calculated rate constants for Mg–O–Si sites in each protonation state were used to calculate the release rate of Mg$^{2+}$ from the forsterite surface according to the model that the surface contains a fraction of sites in each protonation state at every pH and that the number of OH groups available on the surface also contributes to the rate. This model is given by (2-10):

$$Rate = \rho (\theta_{prot} k_{prot} + \theta_{neut} k_{neut} + \theta_{deprot} k_{deprot})$$

where $\rho$ (in units of mol m$^{-2}$) is the reactive site density and thus the number of OH groups on the surface, $\theta_i$ is the fraction of each site on the surface at a given pH, and $k_i$ (in s$^{-1}$) is the rate constant for each $i$ type of site. Thus, the rate of Mg$^{2+}$ release is proportional to the number of OH sites on the surface as well as the sum of the contribution from sites of each protonation state on the surface. The density of reactive
sites, \( \rho \), was extracted from the forsterite (100) surface provided in the Cerius\(^2\) database\(^3\) where the number of OH groups on each Si and each Mg\(^{2+}\) on the surface were totaled. The fraction of protonated, neutral, and deprotonated sites, \( \theta \), was determined from the data of Pokrovsky and Schott where the apparent surface charge was plotted versus pH, and the concentration of H\(^+\) adsorbed to the surface can be obtained from (2-11):

\[
\sigma_{\text{app}} = F[H^\text{ads}]^{-1}
\]

where \( \sigma_{\text{app}} \) is the apparent surface charge, \( F \) is Faraday’s constant, and \( s \) is the surface area of the mineral sample.\(^2\)

Once the amount of H\(^+\) or OH\(^-\) adsorbed to the surface was determined, the number of remaining neutral and deprotonated or protonated sites was estimated. The amount of adsorbed H\(^+\) was treated as the number of protonated Mg–O–Si sites, and the number of deprotonated sites was set at zero according to the pH of zero surface charge of Pokrovsky and Schott.\(^2\) The number of neutral Mg–O–Si sites was determined simply from (2-12):

\[
\theta_{\text{neut}} = 1 - \theta_{\text{prot}} - \theta_{\text{deprot}}
\]

where \( \theta_{\text{neut}} \) is the number of neutral Mg–O–Si sites, \( \theta_{\text{prot}} \) is the number of protonated Mg–O–Si sites, and \( \theta_{\text{deprot}} \) is the number of deprotonated Mg–O–Si sites on the surface at any given pH. These values were then used to calculate the rate of Mg\(^{2+}\) release from the surface according to Equation 2-10.
2.3 Results and Discussion

2.3.1 Hydrolysis of Mg–O–Si Sites

2.3.1.1 Reaction Coordinate

Two possibilities exist for the reaction coordinate of Mg–O–Si hydrolysis reactions: lengthening of the Mg–O bond and decreasing the Mg–O distance between the incoming H₂O molecule and the Mg²⁺. PES scans along the latter reaction coordinate led to H₂O exchange around the Mg²⁺ rather than hydrolysis of the Mg–O bond, suggesting that Mg²⁺ extraction from the surface can be de-coupled from H₂O exchange.

A representative scan for the exchange of H₂O molecules around the Mg²⁺ in the neutral site is shown in Figure 2-4. The reactant complex (RC), transition state (TS), and product complex (PC) structures shown in Figure 2-4a were taken directly from the PES scan, and the energy for each point along the scan is shown in Figure 2-4b. The H₂O molecule from the second hydration sphere is circled in blue, while the H₂O molecule that is originally bonded to the Mg²⁺ and will be replaced is circled in green (Figure 2-4a). In the RC, the H₂O molecule in the second hydration sphere is 3.74 Å from the Mg²⁺, and the H₂O molecule in the first hydration sphere is 2.22 Å from the Mg²⁺. As the H₂O molecule from the second hydration sphere approaches the Mg²⁺, the bond between the Mg²⁺ and the H₂O in the first hydration sphere begins to lengthen. In the TS, this bond length is 2.36 Å, while the H₂O from the second hydration sphere is 2.44 Å from the Mg²⁺. Once these two H₂O molecules have exchanged, the Mg–H₂O bond is 2.14 Å, and the H₂O that is now in the second hydration sphere is 3.49 Å from the Mg²⁺. The energy profile shows that the barrier height for this process is 41 kJ/mol. Although this
Figure 2-4: The PES scan of H₂O exchange around a neutral Mg–O–Si site. (a) The gas phase RC, TS, and PC species from the scan. The H₂O molecule circled in blue represents the H₂O molecule from the second hydration sphere that will bond with the Mg²⁺ in the products, and the H₂O molecule in the green circle will be released from the first hydration sphere of Mg²⁺. (b) The energy (kJ/mol) of each point on the scan is plotted against the Mg–H₂O distance of the incoming H₂O molecule, and thus the progression of the scan is from right to left.
process occurs regularly in solution, the key to understanding dissolution behavior is to determine how the Mg–O bond is broken, resulting in the leaching of Mg\(^{2+}\) species to solution. These scans show that decreasing the H\(_2\)O–Mg distance does not contribute to the release of Mg\(^{2+}\) species from the forsterite surface.

On the other hand, PES scans of the Mg–O distance showed that a change in this parameter does lead to the release of Mg\(^{2+}\) species from the surface. The products for each of these scans of Mg–O bond break contained a penta-coordinated Mg\(^{2+}\) species where the incoming H\(_2\)O molecule had not yet been absorbed. For this reason, a second set of scans was performed to determine how this absorption process occurs. After both sets of scans were completed, further optimization of possible RC, TS, and PC structures from those PES scans led to the isolation of the stationary points on the reaction profile, and both sets of scans are included here.

2.3.1.2 Energy Profiles

The potential energy surface scans and energy profiles for each reaction are presented here for the gas-phase, single-point aqueous-phase, and energy-minimized aqueous-phase. The PES scans represent the energy for the complex at each point along the scan compared to the RC, for which the energy is set as zero. The energy profiles represent the energy of the optimized RC, TS, and PC species, where the RC energy is again set as zero. The PES scans in Figures 2-5, 2-6, and 2-7 were used to gain a more complete understanding of how each reaction occurs, and they represent the two steps in the release of Mg\(^{2+}\) from the surface: the breaking of the Mg–O bond and the absorption of the incoming H\(_2\)O to form a hexa-coordinated product. The barrier heights for these
reactions are taken from the energy profiles and are also shown in Figures 2-5, 2-6, and 2-7, and the reaction with the highest barrier height varies between phases. For the gas-phase calculations, the barrier heights increase in the order of protonated, deprotonated, and neutral site reactions, whereas for the energy-minimized aqueous-phase calculations, the deprotonated site reaction has the lowest barrier height. The barrier heights for the protonated, neutral, and deprotonated sites in each of the three phases are summarized in Table 2-1, and the bond lengths given in the text are those for the gas-phase and are meant to be a guide for the reader.

The PES scan of the Mg–O distance and then the H$_2$O–Mg$^{2+}$ distance for the protonated reaction in the gas-phase is included in Figure 2-5. The estimates for the RC and the TS from this scan are <1 kJ/mol different in energy which shows the structures derived from the scan are good estimates of the optimized structures. However, the PC from the PES scan of the Mg–O distance has a penta-coordinated Mg$^{2+}$ with the incoming H$_2$O H-bonded to the axial H$_2$O molecule and an equatorial H$_2$O molecule. The scan of the H$_2$O–Mg$^{2+}$ distance shows that the H-bonding network opens up to allow the H$_2$O from the second hydration sphere to further approach the Mg$^{2+}$. The PC from the second scan is a hexaaqua Mg$^{2+}$ ion and is lower in energy than the penta-coordinated conformation by 8 kJ/mol. The aqueous-phase PES scans follow the same behavior as the gas-phase.

The gas-phase, single-point aqueous-phase, and energy-minimized aqueous-phase energy profiles for the protonated reaction are also shown in Figure 2-5. The intact Mg–O bond length is 2.17 Å, and as the bond begins to lengthen, the negative frequency in the TS corresponds to the Mg–O bond breakage at 3.07 Å. The bond is fully broken in
Figure 2-5: The energy plot for the protonated Mg–O–Si site reaction. The gas-phase potential energy scan (G-P PES Scan) is green, and the aqueous-phase potential energy scan (A-P PES Scan) is orange. The energy profiles comparing the RC, TS, and PC in the gas-phase (G-P), single-point aqueous-phase (Single-Point A-P), and energy-minimized aqueous-phase (A-P) are black, red, and blue, respectively.

the product complex where the Mg–O distance is 4.14 Å, and the barrier heights for this reaction are 21 kJ/mol in the gas-phase, 33 kJ/mol in the single-point aqueous-phase, and 37 kJ/mol in the energy-minimized aqueous-phase calculations.

The PES scans for the neutral reaction are shown in Figure 2-6. As with the protonated site reaction, the products for the neutral reaction in the scan include a H2O molecule and a penta-coordinated Mg2+. In the PES scan of the H2O–Mg2+ distance, the incoming H2O breaks the H-bond network with the two H2O molecules with which it is interacting, and as with the protonated site, the H2O molecules bonded to Mg2+ rearrange to allow the incoming H2O molecule to bond with the Mg2+. As before, the hexa-coordinated Mg2+ species is lower in energy than the penta-coordinated Mg2+ by more than 20 kJ/mol. This same phenomenon is observed in the gas-phase as well but the difference in energy is 6 kJ/mol.
Figure 2-6: The energy plot for the neutral Mg–O–Si site reaction where the labeling and color scheme are the same as Figure 2-5

The energy profile for the neutral site reaction is given in Figure 2-6, and the gas-phase, single-point aqueous-phase, and energy-minimized aqueous-phase results are included. The initial Mg–O bond distance in the RC is 1.98 Å, shorter than in the protonated state because here the O is not protonated. As the bond lengthens, the breaking of the Mg–O bond in the TS occurs at a Mg–O distance of 2.78 Å. The Mg–O distance continues to lengthen until the PC where it is 3.79 Å, and the barrier heights for this reaction are 54 kJ/mol in the gas-phase, 40 kJ/mol in the single-point aqueous-phase, and 44 kJ/mol in the energy-minimized aqueous-phase calculations, respectively.

The PES scans for the deprotonated site reaction are shown in Figure 2-7. The gas-phase scan mimics the energy values for the RC and TS for this reaction in that they are less than 1 kJ/mol apart for each structure. The aqueous-phase is different in that the
Figure 2-7: The energy plot for the deprotonated Mg–O–Si site reaction where the labeling and color scheme are the same as Figure 2-5.

TS from the PES scan is 7 kJ/mol higher in energy than the energy-minimized gas-phase. In the PES scan of the H₂O–Mg²⁺ distance, the incoming H₂O molecule approaches the Mg²⁺, and this causes the axial OH group to rearrange in order to allow the H₂O to approach. This scan in the aqueous-phase showed decreasing the Mg²⁺–H₂O distance to less than 3.40 Å caused an increase in energy of over 70 kJ/mol, indicating that the penta-coordinated Mg²⁺ with the H₂O molecule in the second hydration sphere is more stable, and a more detailed discussion of the presence of this penta-coordinated Mg²⁺ in the PC will be given in the next section.

The energy profiles for the hydrolysis of deprotonated sites are shown in Figure 2-7 for the gas-phase, single-point aqueous-phase, and energy-minimized aqueous-phase calculations. The initial Mg–O distance is 2.01 Å, and the bond lengthens along the reaction coordinate until breakage in the TS where the Mg–O distance is 2.98 Å. In the
Table 2-1: Barrier heights (kJ/mol) for hydrolysis of protonated, neutral, and deprotonated sites for calculations in the gas-phase, single-point aqueous-phase, and energy-minimized aqueous-phase calculations.

<table>
<thead>
<tr>
<th>Phase</th>
<th>Protonated</th>
<th>Neutral</th>
<th>Deprotonated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas</td>
<td>21</td>
<td>54</td>
<td>39</td>
</tr>
<tr>
<td>Single-Point Aqueous</td>
<td>33</td>
<td>40</td>
<td>21</td>
</tr>
<tr>
<td>Energy-Minimized Aqueous</td>
<td>37</td>
<td>44</td>
<td>40</td>
</tr>
</tbody>
</table>

PC, the Mg–O distance is 3.76 Å, and the barrier heights for this reaction are 39 kJ/mol in the gas-phase, 21 kJ/mol in the single-point aqueous-phase, and 40 kJ/mol in the energy-minimized aqueous-phase.

2.3.1.3 Reaction Mechanisms

In this section, the reaction mechanisms for the hydrolysis of Mg–O–Si sites in protonated, neutral, and deprotonated states are discussed. The reaction mechanisms show that each site releases Mg$^{2+}$ to solution through the breaking of the Mg–O bond, and the Mg$^{2+}$ is hexa-coordinated in the PCs for the protonated and neutral sites but penta-coordinated for the deprotonated site. Depictions of the reaction mechanisms for the hydrolysis of protonated, neutral, and deprotonated sites in the gas-phase appear in Figures 2-8, 2-9, and 2-10, respectively, and the Mg–O and Si–O distances as well as the Mg–O–Si angles for the RCs and TSs for each reaction in the gas-phase are given in Table 2-2.

The reaction mechanism for the hydrolysis of protonated Mg–O–Si sites appears in Figure 2-8. In the RC shown in Figure 2-8, the Mg$^{2+}$ is bonded to five H$_2$O molecules and the O, which in this case is protonated. The protonation of O in conjunction with the
Figure 2-8: The reaction mechanism for the hydrolysis of a protonated Mg–O–Si site. The reactant complex (RC), the transition state (TS), and the product complex (PC) are shown, and the color scheme is the same as Figure 2-1.

Use of H₂O as the Mg–O bond leads to breakage in the TS and leaves Mg²⁺ penta-coordinated. The TS shows attraction of an equatorial H₂O molecule to the O as well as rearrangement of two equatorial H₂O molecules on the Mg²⁺ to accommodate the incoming H₂O molecule. The PC is comprised of a hexaaqua Mg²⁺ ion, showing the incoming H₂O molecule has been absorbed by the Mg²⁺, and silicic acid, and three H₂O molecules on the Mg²⁺ participate in H-bonding with three OH groups on the silicic acid in the PC. This reaction mechanism agrees with previous calculations that showed that Mg²⁺ is released from the surface in the form of Mg(H₂O)₆²⁺ and that the O remains protonated and bonded to the Si atom.⁴⁶

The hydrolysis of neutral Mg–O–Si sites is represented by the reaction of a H₂O molecule with a cluster similar to that pictured in Figure 2-8 but without the protonation of the O, and the mechanism for this reaction appears in Figure 2-9. In the RC, the incoming H₂O molecule is attracted to the axial H₂O molecule and an equatorial H₂O
molecule bonded to the Mg$^{2+}$. The negative frequency in the TS corresponds to a concerted motion where the Mg–O bond is broken at the same time that an H$^+$ from an equatorial H$_2$O molecule is transferred to the O which now remains on the Si species. Also in the TS, breaking of the Mg–O bond in the TS leads to a decrease in Mg$^{2+}$ coordination as with the protonated state. In the PC, the incoming H$_2$O molecule has been absorbed by the Mg$^{2+}$ and also occurred in the protonated site reaction. Hence, the PC is comprised of a hexa-coordinated Mg$^{2+}$, in the form of [Mg(H$_2$O)$_5$OH]$^+$, and silicic acid. The two H$_2$O molecules and the OH group on the Mg$^{2+}$ are engaged in H-bonding with three OH groups on the Si atom.

Lastly, the deprotonated site hydrolysis reaction is depicted in Figure 2-10. The same Mg–O–Si molecular cluster from the neutral site reaction is reacted with OH$^-$ in this case, and this is meant to simulate the existence of a deprotonated site on the surface. At the start of the reaction, an H$^+$ is transferred from the axial H$_2$O group on the Mg$^{2+}$ to the incoming OH$. Thus, in effect, a H$_2$O molecule reacts with the Mg–O–Si cluster.

Figure 2-9: The reaction mechanism for the hydrolysis of neutral Mg–O–Si sites. The color scheme is the same as Figure 2-1, and the labeling is the same as Figure 2-8.
Figure 2-10: The reaction mechanism for the hydrolysis of deprotonated Mg-O-Si sites. The color scheme is the same as Figure 2-1, and the labeling is the same as Figure 2-8.

which now has an axial OH group, and these species are pictured as the RC in Figure 2-10. This reaction of a H$_2$O molecule with the deprotonated site due to a H$^+$ abstraction has been postulated for quartz as well.\textsuperscript{3} In the TS, the Mg–O bond break leaves the Mg$^{2+}$ penta-coordinated. Also in the TS, an equatorial H$_2$O molecule is attracted to the O whose valence is no longer complete, and this causes an H$^+$ to be transferred to the O. The incoming H$_2$O molecule is not, however, bonded to the Mg$^{2+}$ as seen in the previous two reactions, and the distance between the Mg$^{2+}$ and the incoming H$_2$O molecule is approximately 1.0 Å greater than those in the first hydration sphere. Thus, the PC is comprised of silicic acid, a H$_2$O molecule, and [Mg(H$_2$O)$_3$(OH)$_2$].

The PC for this reaction warrants additional comment. In the gas-phase, a penta-coordinated PC is expected because the number of negatively charged groups attached to Mg$^{2+}$ causes a decrease in its coordination number,\textsuperscript{47,48} and this PC is logical because the two OH groups can donate excess electrons to the Mg$^{2+}$ within a small volume.\textsuperscript{49} For completeness, a hexa-coordinated version of the PC was optimized in both the gas- and
aqueous-phases, and this hexa-coordinated structure was higher in energy than the penta-coordinated form with the incoming H₂O molecule in the second hydration sphere by more than 20 kJ/mol in each phase. Furthermore, aqueous Mg²⁺ species can also exist in lower coordinated states⁴⁸-⁵⁰ which depend upon the nature of the bonding groups.⁴⁷-⁴⁹ The possibility of producing either Si(OH)₄ or SiO(OH)₃⁻, representing the hydroxylated and deprotonated surface, respectively, as products will be discussed below.

The transfer of the H⁺ to the O in the gas-phase deprotonated site reaction is anomalous because the pKₐ values of aqueous Mg²⁺ species are higher than those for Si⁵¹,⁵² which would indicate that H⁺ transfer is unlikely for these systems, and therefore, SiO(OH)₃⁻ rather than Si(OH)₄ would be the expected Si species in the products at high pH. In the gas-phase PES scan for the deprotonated site, the proton transfer occurs alongside of the Mg–O bond break; that is, as the Mg–O distance increases, the H–O distance decreases. The final bond length of the H⁺ transferred to the O residing on the Si species is 1.05 Å.

This anomaly prompted the use of aqueous-phase PES scans with small steps in the Mg–O distance to isolate the point in the scan at which the proton transfer occurred. In fact, for the PES scans in the aqueous-phase, the opposite is true. In both the neutral and deprotonated site reactions, the H⁺ remains on the original O atom, albeit with elongated bonds of 1.04 and 1.03 Å in the neutral and deprotonated site reactions, respectively. Thus, the results described here in the aqueous-phase show that the Si species in the PC is SiO(OH)₃⁻ and is the immediate product from the hydrolysis reaction.
The Mg–O and Si–O distances as well as the Mg–O–Si bond angles for RCs and TSs in the gas-phase for all reactions are given in Table 2-2. The Mg–O and Si–O distances for the RC in the protonated site reaction reflect that the O is protonated as exhibited by the longer bond for this protonation state. The Mg–O distance in the transition states is longest for the protonated state, followed by the deprotonated and neutral states. The Mg–O distance in the deprotonated state TS is longer than in the neutral site as a result of donated electron density from the axial OH group on the Mg$^{2+}$. This additional electron density throughout the cluster stabilizes the Mg–O distance in the TS and allows the distance to be larger than in the neutral state TS. Also in the TS, the Si–O distances for the neutral and deprotonated state reactions are shorter than that for the protonated state because the proton has not yet fully attached to the O, and thus excess electron density surrounds the O in the Si–O bond.

Lastly, the Mg–O–Si angles are similar for each protonation state in the RCs in that they differ by only 4%. This shows that the protonation of O does not have an effect on this angle, but the fact that this angle is over 120° represents compensation for steric crowding within this cluster. In addition, the beginning of a tetrahedral geometry around the O becomes evident in each of the transition states as the angles are 110° – 112°.

2.3.1.4 Gibbs Free Energy Change

In addition to the reaction mechanisms and barrier heights, the change in Gibbs free energy (ΔG) for each reaction in all three phases is presented in Table 2-3. The overall trend is that these reactions are spontaneous at room temperature, and this is evident particularly in the calculations for the single-point and energy-minimized
Table 2-2: Mg–O and Si–O distances (Å) and Mg–O–Si angles (°) for RCs and TSs for all reactions in the gas-phase.

<table>
<thead>
<tr>
<th>Protonation State</th>
<th>Stationary Point</th>
<th>Mg–O (Å)</th>
<th>Si–O (Å)</th>
<th>Mg–O–Si Angle (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Protonated</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RC</td>
<td>2.17</td>
<td>1.70</td>
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<td>129</td>
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<tr>
<td>TS</td>
<td>3.07</td>
<td>1.68</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>RC</td>
<td>1.98</td>
<td>1.59</td>
<td></td>
<td>127</td>
</tr>
<tr>
<td>TS</td>
<td>2.78</td>
<td>1.58</td>
<td></td>
<td>110</td>
</tr>
<tr>
<td><strong>Deprotonated</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RC</td>
<td>2.01</td>
<td>1.59</td>
<td></td>
<td>126</td>
</tr>
<tr>
<td>TS</td>
<td>2.98</td>
<td>1.58</td>
<td></td>
<td>110</td>
</tr>
</tbody>
</table>
Table 2-3: The change in Gibbs free energy (ΔG) values (kJ/mol) for Mg\(^{2+}\) release from protonated, neutral, and deprotonated sites in the gas- and aqueous-phase hydrolysis reactions.

<table>
<thead>
<tr>
<th>Site</th>
<th>Gas-Phase</th>
<th>Single-Point Aqueous-Phase</th>
<th>Energy-Minimized Aqueous-Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protonated</td>
<td>–2</td>
<td>–14</td>
<td>–12</td>
</tr>
<tr>
<td>Neutral</td>
<td>38</td>
<td>13</td>
<td>–8</td>
</tr>
<tr>
<td>Deprotonated</td>
<td>23</td>
<td>–2</td>
<td>–29</td>
</tr>
</tbody>
</table>

aqueous-phase reactions. The neutral and deprotonated site reactions are not thermodynamically favorable in the gas-phase but spontaneous in the energy-minimized aqueous-phase calculations. This suggests that the ΔG from gas-phase calculations is misleading. Therefore, determination of ΔG values for geochemical reactions investigated via ab-initio methods should be performed using energy-minimized aqueous-phase calculations as opposed to those in the gas-phase.

2.3.1.5 Overall Trends and Comparison to Experiment

Each of the hydrolysis reactions for Mg–O–Si sites proceeds through a dissociative interchange (I\(_d\)) mechanism.\(^{53}\) The I\(_d\) mechanism is manifested by the decrease in coordination of the Mg\(^{2+}\) between the RC and the TS caused by the breaking of the Mg–O bond, and the Mg\(^{2+}\) returns to hexa-coordination in the PC for both the protonated and neutral states after the incoming H\(_2\)O molecule bonds with the Mg\(^{2+}\). Further, the decrease in coordination is expected in the TS because Mg\(^{2+}\) cannot accommodate seven bonds in its first hydration sphere.\(^{50}\) Although the Mg\(^{2+}\) is penta-
coordinated in the PC for the deprotonated site reaction, another H$_2$O molecule is likely to be bonded with the Mg$^{2+}$ when this reaction occurs in the presence of bulk water.$^{53}$

The calculated Mg–O–Si barrier heights given in Table 2-1 can be compared with previous calculations of barrier heights for Si–O–Si site hydrolysis,$^{2,24,25,54}$ and these values appear in Table 2-4. The barrier heights in the gas-phase for Mg–O–Si site hydrolysis for each protonation state are lower than those for Si–O–Si hydrolysis in the same protonation state. Therefore, the Mg–O–Si barrier heights for Mg$^{2+}$ release presented here in conjunction with the Si–O–Si barrier heights in Table 2-4 indicate that the Mg$^{2+}$ species would be released from the forsterite surface first. Further, this comparison can be extended to include the barrier heights for other silicate minerals as well. The barrier heights in Table 2-1 for Mg–O–Si site hydrolysis are also lower than those for the hydrolysis of Si–O–Al sites in the same protonation state as well as for Al–O–Si sites in the pH range at which each site would exist.$^4$ When these Al-bearing sites are in turn compared with the values in Table 2-4 for Si–O–Si site hydrolysis, the Si–O–Si sites have the highest barrier heights. When one considers this particular group – Mg$^{2+}$, Al$^{3+}$, and Si$^{4+}$ – the strength of the M–O bond increases as one moves to higher atomic number across the row in the periodic table when the octets of Mg$^{2+}$ and Al$^{3+}$ are expanded. This trend correlates with the observation that dissolution rates in the same pH range decrease for Mg$^{2+}$ to Al$^{3+}$ to Si$^{4+}$ silicate minerals.$^{21}$

Several previous experimental descriptions of forsterite dissolution have included measurements of the activation energy ($E_a$). However, these are actually “apparent activation energies ($E_{app}$).”$^{55-58}$ The dissolution of forsterite includes the leaching of both Mg$^{2+}$ and Si from the surface and from sites in all protonation states as well as the
Table 2-4: Ab-initio barrier heights (kJ/mol) for Si–O–Si site hydrolysis in the gas phase.\textsuperscript{2,24,25,54}

<table>
<thead>
<tr>
<th>Source</th>
<th>Method</th>
<th>Protonated</th>
<th>Neutral</th>
<th>Deprotonated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xiao and Lasaga 1994</td>
<td>HF/6-31G(d), MP2/6-31G(d)</td>
<td>100, 92</td>
<td>121, 134</td>
<td></td>
</tr>
<tr>
<td>Xiao and Lasaga 1996</td>
<td>MP2/6-31G(d)</td>
<td></td>
<td></td>
<td>79</td>
</tr>
<tr>
<td>Pelmenschikov \textit{et al.}</td>
<td>B3LYP/6-31G(d), B3LYP/6-311G(d,p)</td>
<td></td>
<td>105</td>
<td></td>
</tr>
<tr>
<td>Nangia and Garrison</td>
<td>B3LYP/6-31+G(d,p), B3LYP/MG3S</td>
<td>69, 75</td>
<td>159, 174</td>
<td>110, 122</td>
</tr>
</tbody>
</table>
transport of $\text{H}_3\text{O}^+$, $\text{H}_2\text{O}$, and $\text{OH}^-$ species to the surface,\textsuperscript{55} their adsorption,\textsuperscript{56} and overall network opening. Furthermore, this $E_{\text{app}}$ term also includes contributions from temperature-dependent processes that affect dissolution\textsuperscript{55,57} such as surface speciation.\textsuperscript{57} Moreover, if the experiment has not reached steady state, this can have a profound effect on $E_{\text{app}}$.\textsuperscript{58} Furthermore, the scatter in previous data depends mainly on the temperature at which the experiment was performed. The temperature is important because experimentally measured $E_{\text{app}}$ values are extracted from the slope of the line formed when the log of the dissolution rate constant ($k$) is plotted versus the reciprocal of the temperature (in K). At temperatures below 327K, those processes in dissolution that have low $E_a$ barriers dominate, and at temperatures higher than 327K, high $E_a$ processes dominate.\textsuperscript{59} Thus, the $E_{\text{app}}$ that is measured in a dissolution experiment hinges upon the temperature at which the experiment was performed, and these aspects of dissolution must be considered when an $E_{\text{app}}$ is communicated.

Previous researchers have calculated experimental $E_{\text{app}}$ values for the dissolution of forsterite. Van Herk \textit{et al.} performed dissolution experiments from pH = 1 to 3 over a temperature range of 313 to 343K, and they determined $E_{\text{app}}$ values of 25 ± 2 kJ/mol using hydrochloric acid and 30 ± 2 kJ/mol using sulfuric acid.\textsuperscript{60} Awad \textit{et al.} also investigated the acidic dissolution of forsterite, specifically at pH = 1 to 2 and over the temperature range 296 to 363K. They determined the $E_{\text{app}}$ for the dissolution of the bulk crystal to be 72 ± 12 kJ/mol.\textsuperscript{20} The difference between these two values has been attributed to the inherent physical nature of the forsterite samples used.\textsuperscript{20} In addition, the discrepancies between the results presented in this work shown in Table 2-1 and these experimental values reflect the fact that these values represent different processes.
Moreover, the protonated Mg–O–Si site barrier heights are lower than nearly all of the values calculated by Van Herk et al. and Awad et al., and this represents the fact that breaking the Mg–O bond is not the rate-limiting step.

However, there are experimental data that support the hypothesis that forsterite dissolution is controlled by Si–O–Si hydrolysis rather than Mg–O–Si hydrolysis. Pokrovsky and Schott performed dissolution experiments at pH \( \leq 12 \) and observed the formation of a Si-rich, Mg\(^{2+}\)-depleted layer on their forsterite samples at pH < 9,\(^{23}\) and they postulated that the layer was comprised of polymerized silica tetrahedra.\(^{8}\) Forsterite dissolution in acidic solution was investigated via \(^{29}\)Si MAS NMR by Davis and coworkers using a gem quality sample with no known impurities,\(^{61}\) and their primary finding was that Mg\(^{2+}\) was released from the surface first, leaving behind a Si-rich layer. The forsterite was dissolved at an initial pH of 3, and as Mg\(^{2+}\) ions were replaced by two H\(^{+}\) ions at each site, the pH increased accordingly. This led to the development of a silica-type layer on the surface that was depleted of Mg\(^{2+}\). The \(^{29}\)Si MAS NMR spectra from these experiments showed the development of polymerized Si species in the form of Q\(^3\) Si–O–Si units, or Si atoms connected through O atoms to three other Si atoms, which increased in concentration over time. The original forsterite sample was comprised of Q\(^0\) Si units – that is, those without Si–O–Si units – which after dissolution give way to a Si-rich layer whose structure was composed of Si–O–Si bonds. Therefore, this recent experimental description of forsterite dissolution shows that Mg\(^{2+}\) release from the surface leads to the development of a Si-rich layer which is comprised of Q\(^3\) sites and is in contrast to the original forsterite mineral structure that is characterized by Q\(^0\) silicon sites.
2.3.2 Mg$^{2+}$ Release Rates from Mg–O–Si Sites versus pH

2.3.2.1 Parameters for Calculating the Rate

The pre-exponential factors and rate constants for the hydrolysis reactions at protonated, neutral, and deprotonated sites in the gas-, single-point aqueous-, and energy-minimized aqueous-phases are given in Table 2-5, and the rate constants follow the same trend as the barrier heights outlined above. The rates for the release of Mg$^{2+}$ from forsterite were calculated using the model from Equation 2-10 and appear in Table 2-6. The data in Table 2-6 are calculated for pH = 3.0 to 11.2, and rates for gas-, single-point aqueous-, and energy-minimized aqueous-phases are grouped in that order. The calculated rates for Mg$^{2+}$ release given in Table 2-6 decrease with increasing pH in the gas-phase and energy-minimized aqueous-phase calculations, and the trend in the single-point aqueous-phase calculations is anomalous due to the barrier heights from these calculations. Although the rate constants inserted into Equation 2-14 were calculated in this work, the $\rho$ and $\theta_i$ values were extracted from experimental data. Twenty-three OH groups appear on 1 nm$^2$ of the forsterite (100) surface which corresponds to a reactive site density of $3.82 \times 10^{-5}$ mol m$^{-2}$. The $\theta_{\text{prot}}$ and $\theta_{\text{deprot}}$ values were taken from Pokrovsky and Schott, and the $\theta_{\text{neut}}$ values were calculated according to Equation 2-12. The elementary rate constants, $k_i$, provided in Table 2-5, for the gas-, single-point aqueous-, and energy-minimized aqueous-phase reactions, were combined with the same
Table 2-5: Pre-exponential factors $A$ (s$^{-1}$) and rate constants $k$ (s$^{-1}$) for the hydrolysis of Mg–O–Si sites on the forsterite surface.

<table>
<thead>
<tr>
<th>Site</th>
<th>Gas-Phase</th>
<th>Single-Point Aqueous-Phase</th>
<th>Energy-Minimized Aqueous-Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$A$</td>
<td>$k$</td>
<td>$A$</td>
</tr>
<tr>
<td>Protonated</td>
<td>$2.4 \times 10^{12}$</td>
<td>$5.7 \times 10^8$</td>
<td>$2.4 \times 10^{12}$</td>
</tr>
<tr>
<td>Neutral</td>
<td>$7.7 \times 10^{11}$</td>
<td>$2.7 \times 10^4$</td>
<td>$7.7 \times 10^{11}$</td>
</tr>
<tr>
<td>Deprotonated</td>
<td>$7.1 \times 10^{11}$</td>
<td>$2.2 \times 10^6$</td>
<td>$7.1 \times 10^{11}$</td>
</tr>
</tbody>
</table>
Table 2-6: Reaction rates (mol m$^{-2}$ s$^{-1}$) for the hydrolysis of protonated, neutral, and deprotonated sites in gas-, single-point aqueous-, and energy-minimized aqueous-phases.

<table>
<thead>
<tr>
<th>pH</th>
<th>$\rho$ (mol m$^{-2}$)</th>
<th>$\theta_{prot}$ (s$^{-1}$)</th>
<th>Protonated (s$^{-1}$)</th>
<th>$\theta_{neut}$ (s$^{-1}$)</th>
<th>Total Rate (mol m$^{-2}$ s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.0</td>
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</tr>
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<td>1.6x10$^{-5}$</td>
<td>5.7x10$^8$</td>
<td>2.3x10$^{-5}$</td>
<td>3.38x10$^{-1}$</td>
</tr>
<tr>
<td>4.7</td>
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<td>1.3x10$^{-5}$</td>
<td>5.7x10$^8$</td>
<td>2.6x10$^{-5}$</td>
<td>2.73x10$^{-1}$</td>
</tr>
<tr>
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<td>8.6x10$^{-6}$</td>
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<td>3.0x10$^{-5}$</td>
<td>1.87x10$^{-1}$</td>
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<td>8.1x10$^{-6}$</td>
<td>5.7x10$^8$</td>
<td>3.0x10$^{-5}$</td>
<td>1.77x10$^{-1}$</td>
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</tr>
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</tr>
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</tr>
<tr>
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<td>$\theta_{\text{prot}}$ (s$^{-1}$)</td>
<td>Protonated (s$^{-1}$)</td>
<td>$\theta_{\text{neut}}$ (s$^{-1}$)</td>
<td>Neutral (s$^{-1}$)</td>
</tr>
<tr>
<td>-----</td>
<td>---------------------</td>
<td>------------------------------</td>
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<td>------------------</td>
</tr>
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<td>3.0x10$^{4}$</td>
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<tr>
<td>11.2</td>
<td>3.82x10$^{-5}$</td>
<td>0.0x10$^{0}$</td>
<td>1.7x10$^{6}$</td>
<td>3.5x10$^{-5}$</td>
<td>3.0x10$^{4}$</td>
</tr>
</tbody>
</table>
\( \rho \) and \( \theta_i \) values in Equation 2-10 to yield the Mg\(^{2+} \) release rate in units of mol m\(^{-2} \) s\(^{-1} \) as given in the last column of Table 2-6.

2.3.2.2 Rates versus pH

In both the gas-phase and energy-minimized aqueous-phase calculations, the rate decreases as pH increases over the range of pH from 3.0 – 11.2. The Mg\(^{2+} \) release rate from the single-point aqueous-phase calculations shows a somewhat U-shaped behavior, which is a result of the anomaly in barrier heights described in Section 2.3.1.2. The need for energy-minimized aqueous-phase calculations for these systems is reinforced here with the anomalous behavior of the single-point aqueous-phase calculations.

2.3.2.3 Comparisons with Previous Work

The approach used here for the rate equation describing the release of species as a function of the types of sites on a surface has been used for quartz, and the calculated rates derived from ab-initio barrier heights replicated the findings of previous quartz dissolution experiments.\(^2\) However, several different parameters are included here that were not present in the previous work. The number of OH groups available for reaction on a forsterite surface has not been measured experimentally, so an estimate was made based on the database included in the Cerius\(^2\) software.\(^30\) Additionally, the experimental data of Pokrovsky and Schott\(^23\) were used to describe the fraction of protonated, neutral, and deprotonated sites on the surface as opposed to detailed descriptions of protonation
state fractions on the quartz surface. The subsequent extrapolation of the number of neutral and deprotonated sites in acidic pH and the number of protonated and neutral sites in basic pH is thus a blend of experimental data and data fitting in this model.

In addition to differences in the methodology of the quartz dissolution rate analysis versus this work, the dissolution chemistry is different between these two minerals. For quartz, overall dissolution occurs as a result of the network opening, site protonation, Si release from Si–O–Si sites, and the net release of material that results from excess dissolution versus precipitation. For forsterite, comparable processes occur for both Mg–O–Si and Si–O–Si sites, which form after the leaching of Mg$^{2+}$ from the surface. Therefore, the discrepancies that exist for the dissolution rates stem from the fact that the Mg$^{2+}$ leaching process is not rate-limiting in the pH ranges studied here, but the replication of experimental trends shows that the model is appropriate for describing Mg$^{2+}$ release.

Stoichiometric dissolution is seen for forsterite dissolution in acidic and basic pH despite the slower release rate of Si to solution and occurs when the surface species are released to solution such that a ratio of their rates is equal to their ratio in the parent mineral. The presence of stoichiometric dissolution in forsterite experiments is intriguing in that Mg$^{2+}$ release results from a lower energy process – that is, the breaking of the Mg–O bond – versus the same process for the release of Si to solution. However, stoichiometric dissolution for this mineral is explained by the sequential release of Mg$^{2+}$ then Si from the forsterite surface. Once the Mg$^{2+}$ is depleted in the surface layer, Si atoms are released from the polymerized layer that develops. Dissolution experiments performed on acid-reacted forsterite samples showed that Si was released preferentially.
indicative of the need to dissolve away this Si-rich layer before additional Mg\(^{2+}\) can be released.

2.4 Conclusions

The mechanisms and rate of Mg\(^{2+}\) release from forsterite have been successfully modeled by integrating experimental data and ab-initio results. The reaction mechanisms and barrier heights of the hydrolysis of Mg–O–Si sites were investigated, and the hydrolysis of the protonated, neutral, and deprotonated states proceeds through a two-step mechanism where the breaking of the Mg–O bond corresponds to the step which releases Mg\(^{2+}\) to solution. The barrier heights for the energy-minimized gas-phase and aqueous-phase calculations show that the protonated sites are the lowest, followed by the deprotonated and neutral sites in turn. Furthermore, any aqueous-phase calculations performed to represent geochemical processes must be optimized as the single-point aqueous-phase calculations do not always yield reliable results.

The barrier heights for the release of Mg\(^{2+}\) from Mg–O–Si sites were used to calculate rate constants and in turn a rate for this process. The rate constants and rates are higher than experimental rates for overall dissolution\(^5\text{--}^{11}\) as indicated by the lower calculated barrier heights compared to \(E_{\text{app}}\) values for the dissolution for forsterite;\(^{20,60}\) thus, the release of Mg\(^{2+}\) is not the rate-limiting step. However, these ab-initio results corroborate previous experimental and ab-initio results;\(^{29}\)\(^{\text{Si MAS NMR experiments of forsterite dissolution show the Mg}}^{2+} \text{is released before Si and that a Si-rich layer develops on the surface.}^{61}\) In conjunction with previous ab-initio calculations for the hydrolysis of
Si–O–Si sites, the ab-initio results here also show that Mg$^{2+}$ leaches first from the forsterite surface.

Initially, two possibilities existed for the reaction coordinate of these reactions, but the decrease in the distance between the H$_2$O molecule in the second hydration sphere and the Mg$^{2+}$ in the Mg–O–Si cluster did not lead to a breaking of the Mg–O bond. Instead, PES scans indicate that the distance between the H$_2$O molecule and the Mg$^{2+}$ led only to an exchange of H$_2$O molecules around the Mg$^{2+}$ when it was hexa-coordinated, whereas the Mg–O bond remained intact. Therefore, the release of Mg$^{2+}$ from the surface as a result of breaking the Mg–O bond is a direct result of the lengthening of the Mg–O bond and not the exchange of H$_2$O molecules around the Mg$^{2+}$. 
2.5 References

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Chapter 3

Quantum Mechanical Modeling of Hydrolysis and H₂O-Exchange in Mg-, Ca-, and Ni-Silicate Clusters: Implications for Dissolution Mechanisms of Olivine Minerals

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3.1 Introduction

The work described in Chapter 3 employs quantum mechanical calculations to investigate the proposed correlation between H₂O exchange rates around aqueous metal ions and the dissolution rates of silicate minerals containing those ions. The H₂O exchange reaction for a hydrated Mg²⁺ ion is modeled in order to determine whether gas-phase or aqueous-phase approximations are more appropriate to model these systems. The conditions that cause each M–O bond to break, namely H₂O approach of M–O bond lengthening, are identified. The barrier heights for H₂O exchange and hydrolysis reactions for each protonated M–O–Si (M = Ni²⁺, Mg²⁺, Ca²⁺) site are calculated and used to calculate the rate constant for each of these reactions. For all of the reactions studied, a comparison with experimental observations is made.

Dissolution occurs via reaction of species from solution with sites on the mineral surface, leading ultimately to the breakdown of the original mineral structure and resulting in the release of elemental constituents to solution, as in **Reactions 3-1 and 3-2**.
Depending upon whether the mineral is forsterite or anorthite, dissolution of the mineral in acidic media would proceed as \((3-1)\):\(^2\,^5\)

\[
\text{Mg}_2\text{SiO}_4(\text{aq}) + 4\text{H}^+(\text{aq}) \rightarrow 2\text{Mg}^{2+}(\text{aq}) + \text{Si(OH)}_4(\text{aq})
\]

for forsterite or \((3-2)\):

\[
\text{CaAl}_2\text{Si}_2\text{O}_8(\text{aq}) + 2\text{H}^+(\text{aq}) + \text{H}_2\text{O}(\text{l}) \rightarrow \text{Ca}^{2+}(\text{aq}) + \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4(\text{s})
\]

for anorthite. This process is comprised of many reactions, including the opening of the surface structure network, the replacement of the O atoms in the metal surface polyhedra with H\(_2\)O molecules,\(^10\) and surface site hydrolysis. This last class of reactions occurs when a H\(_3\)O\(^+\) ion, H\(_2\)O molecule, or OH\(^-\) ion from solution reacts with a metal site on the surface to release a metal ion to solution, which results from the breaking of the M–O bond.

The H\(_2\)O molecules, H\(_3\)O\(^+\) ions, and OH\(^-\) ions from solution affect the dissolution rate through two possible mechanisms. The effect of the H\(_3\)O\(^+\) ion on the dissolution rate has been described as proton-promoted dissolution,\(^11,\,12\) where the dissolution rate of a mineral is observed to increase as a function of decreasing pH. This is likely because H\(^+\) ions adsorb to surface O atoms and weaken the M–O and Si–O bonds. The second proposed mechanism is known as ligand-promoted dissolution.\(^11-\,13\) The H\(_2\)O molecule and OH\(^-\) ion serve as ligands bonding to the mineral surface, and their effect on the dissolution rate likely results from the weakening of the M–O surface bonds via an increase in charge to the metal that is delivered by the adsorbing ligand.\(^11\) At a given pH, there is a distribution of H\(^+\), H\(_2\)O, and OH\(^-\) species from solution reacting with surface
sites, and thus the proton- and ligand-promoted pathways can both be active during dissolution.12

While ligand-promoted dissolution occurs as a result of the exchange of ligands from solution for H2O molecules or OH– groups on the surface,13 a ligand exchange reaction occurs when a group on an aqueous metal species is replaced by another from solution, and this reaction proceeds as (3-3):

$$MX_n + Y \leftrightarrow MX_{n-1}Y + X$$

where M is an aqueous metal ion, X is a ligand coordinated with M, and Y is a ligand from solution, according to the notation of Langford and Gray.14 One type of ligand exchange reaction is H2O exchange around metal ions in solution.15 Such reactions can also occur on a mineral surface, and in the case of silicate minerals, this process involves the breaking and forming of M–O bonds at surface sites. Rates of H2O exchange reactions around a metal cation in solution correlate with the amount of energy required to break the M–O bond15,16 as well as with the dissolution rates of orthosilicate minerals.15,17-19 This trend arises because both H2O exchange reactions and the release of cations from silicate minerals involve the breaking of M–O bonds – in the form of M–O, M–OH, or M–OH2, and this correlation extends across nearly seven orders of magnitude in orthosilicate dissolution rates.17 Because both processes involve breaking of M–O bonds, the reaction mechanisms for these two processes have been hypothesized to be fundamentally similar,13,17 and previous researchers have suggested that this similarity enables a direct comparison between rates for ligand-promoted dissolution and H2O exchange reactions in solution.13 However, the amount of energy required to break a M–
O bond decreases across the following series: M–O > M–OH > M–OH₂. Therefore, the rates of such reactions are not equal, and the correlation of dissolution rates of orthosilicate minerals with H₂O exchange rates may not mean that the mechanisms are the same.

Both H₂O exchange reactions and ligand-promoted dissolution reactions in the form of hydrolysis of M–O bonds occur at surface sites during dissolution of silicate minerals. What remains to be seen is if these two phenomena are related and if the H₂O exchange rate of an aqueous metal ion is an indicator of how dissolution proceeds for a mineral containing that cation. There is no known chemical explanation¹⁵ for the correlation of H₂O exchange reaction rates and the dissolution rates of orthosilicate minerals.¹⁵,¹⁷⁻¹⁹ H₂O molecules are thought to replace O atoms in mineral polyhedra during dissolution in acidic pH.¹⁰ Although extensive experimental data exist,¹⁰,¹⁵,¹⁷,²⁰,²¹ molecular mechanisms of these surface reactions have yet to be described for divalent silicate minerals using quantum mechanical techniques. The link between H₂O exchange reactions in solution and the dissolution of divalent silicate minerals has not been investigated from a molecular perspective. Our hypothesis is that the release of metals to solution occurs as a result of hydrolysis of M–O linkages in the mineral network. One aim of this work is to test the hypothesis that H₂O exchange reactions on the surface lead to M–O bond hydrolysis.

One approach for investigating the link between H₂O exchange reactions and the dissolution rates of minerals is to use quantum mechanical calculations, which enable a molecular scale investigation into reaction mechanisms. Previous work has studied the hydrolysis of sites on the surface for quartz,²² aluminosilicates,²³⁻²⁵ and forsterite.²⁶
These studies analyzed experimental phenomena by using molecular clusters to simulate sites on the mineral surface, and experimental observations were replicated. What is most important about this approach is that the most fundamental aspect of a chemical phenomenon is investigated without the interference of atoms that do not participate in the reaction. Therefore, a molecular cluster investigation would allow for the analysis of H$_2$O exchange and hydrolysis reactions on the surface, enable a description of the reaction mechanisms, and provide insight into the link between H$_2$O exchange reaction rates for metals in solution and the dissolution rates for minerals containing these metals.

Questions that have yet to be answered include whether H$_2$O exchange reactions are coupled with dissolution, how H$_2$O exchange reactions affect dissolution, and if the metal cations studied are released from the mineral surface with similar mechanisms. The results from the quantum mechanical calculations employed in this work seek to answer these questions. Molecular clusters representing protonated M–O–Si (M = Ni$^{2+}$, Mg$^{2+}$, and Ca$^{2+}$) sites on a mineral surface were reacted with H$_3$O$^+$ ions to examine H$_2$O exchange and hydrolysis reactions and to determine whether these processes are coupled for the end-member silicate minerals chosen. These particular ions were chosen for this study because they represent orthosilicate minerals with nearly the slowest through the fastest dissolution rates in recent experimental investigations. This work includes the calculation of activation energy barrier heights and rate constants for these reactions, a description of each reaction mechanism, and a comparison with experimental results.
3.2 Computational Methods

Molecular clusters are employed to model two classes of reactions in this work. The first is a H₂O exchange reaction around the hexaaqua Mg²⁺ ion, Mg(H₂O)₆²⁺, and the second models reactions with H₂O and molecular clusters representing M–O–Si sites on olivine surfaces. In the H₂O exchange reaction around the Mg(H₂O)₆²⁺ ion, Mg(H₂O)₆²⁺ reacts with a H₂O molecule to model the replacement of a H₂O molecule from the first hydration sphere by one from the second. This reaction proceeds as (3-4):

\[
[Mg(H₂O)₅(H₂O)₁st]²⁺ + H₂O₂nd → [Mg(H₂O)₅(H₂O)₂nd]²⁺ + H₂O₁st
\]

with the H₂O molecule originally in the first hydration sphere now in the second.

In the second class of reactions, molecular clusters are employed to represent protonated M–O–Si sites on olivine surfaces. A metal ion – where the metal is Ni, Mg, or Ca – is hexa-coordinated via five H₂O molecules and the M–O–Si linkage. The Si is bonded to three OH groups to represent the bonds to the bulk crystal. This termination via OH groups is supported by the fact that atoms farther than the next nearest neighbor position participate minimally in bond breaking at the surface. These molecular clusters are each reacted with a H₂O molecule to investigate H₂O exchange and hydrolysis reactions occurring at protonated M–O–Si surface sites, where protonation is represented by a H⁺ bonded to the O in the M–O–Si linkage. The presence of both the H⁺ and the reacting H₂O molecule represent the reaction of a H₃O⁺ ion from solution with a M–O–Si site on the mineral surface.
The reactions of these protonated clusters with a H$_2$O molecule are given in Reactions 3-5 through 3-9. The protonated Ni–O–Si site is reacted with a H$_2$O molecule according to (3-5):

\[
[Ni(H_2O)_5(OH)Si(OH)_3]^{2+} + H_2O \rightarrow [Ni(H_2O)_6]^{2+} + Si(OH)_4
\]

forming a hexaaqua Ni$^{2+}$ ion and silicic acid as products.

For protonated Mg–O–Si sites, two reactions are investigated. The first is a H$_2$O exchange reaction (3-6),

\[
[Mg(H_2O)_4(H_2O)_{1st}(OH)Si(OH)_3]^{2+} + H_2O_{2nd} \rightarrow [Mg(H_2O)_4(H_2O)_{2nd}(OH)Si(OH)_3]^{2+} + H_2O_{1st}
\]

where H$_2$O$_{1st}$ is the H$_2$O molecule from the first hydration sphere that moves to the second, and H$_2$O$_{2nd}$ is the H$_2$O molecule from the second hydration sphere that moves to the first. The second Mg–O–Si reaction studied is the hydrolysis of a protonated Mg–O–Si site (3-7),

\[
[Mg(H_2O)_5(OH)Si(OH)_3]^{2+} + H_2O \rightarrow [Mg(H_2O)_6]^{2+} + Si(OH)_4
\]

and the hexaaqua Mg$^{2+}$ ion and silicic acid form as the products.

For protonated Ca–O–Si sites, two reactions are investigated, and the first is H$_2$O exchange (3-8),

\[
[Ca(H_2O)_5(OH)Si(OH)_3]^{2+} + H_2O \rightarrow [Ca(H_2O)_6(OH)Si(OH)_3]^{2+}
\]

where instead of an exchange of H$_2$O molecules from the first and second hydration spheres, the absorption of the second hydration sphere H$_2$O molecule into the primary hydration sphere forms a hepta-coordinated Ca$^{2+}$. The hydrolysis of a protonated Ca–O–Si site proceeds as (3-9):
and the hexaaqua Ca\(^{2+}\) ion and silicic acid form in the products.

The silicic acid product, Si(OH)_4, in Reactions 3-5, 3-7, and 3-9 represents a hydroxylated Si site that remains on the surface after release of the metal to solution. The existence of charge is included in the calculations by assigning the overall system a charge of +2. Thus this charge is dispersed throughout all the atoms in the system and is not solely located on the metal ion. However, for clarity in this work as well as for ease in comparison with experimental data, the metals will each be referred to in their ionic state.

Density functional theory (DFT) calculations have been shown to describe surface site hydrolysis reactions successfully. In particular, the B3LYP functional is used because of its hybrid density functional nature that combines exchange-correlation, gradient correction, and a percentage of Hartree-Fock exchange. The 6-31G(d,p) basis set has been used to model the hydration of Mg\(^{2+}\) and here is used to investigate the reactions in Reactions 3-4 through 3-9. The absence of diffuse functions is advantageous for decreasing the basis set superposition error (BSSE) on transition metal complexes; BSSE is expected to be minimal as we report relative energy changes for molecular clusters, where the intermolecular distances do not vary dramatically.

Molecular clusters representing Mg(H_2O)_6\(^{2+}\) or protonated M–O–Si (M = Ni, Mg, and Ca) sites were optimized with a H_2O molecule in the second hydration sphere, and this optimized structure served as the starting reactants for potential energy surface (PES) scans of the H_2O exchange and hydrolysis reactions. For the H_2O exchange reactions, the
distance between the H$_2$O in the second hydration sphere and the metal was constrained and decreased by 0.1 Å for 20 steps to represent the approach of a H$_2$O molecule from solution. This reaction coordinate was chosen as opposed to varying the distance between the metal and the first hydration sphere H$_2$O molecule because the latter underestimates bound water lifetimes$^{34}$ and because the second hydration sphere H$_2$O molecule distance from the metal is considered fundamental to the H$_2$O exchange reaction.$^{35}$ If the M–O bond did not break during the H$_2$O exchange reaction, then a scan of the M–O distance was performed until it broke, and a second scan of the approach of the second hydration sphere H$_2$O molecule was performed. Each of these scans was comprised of 20 steps, where each step was 0.1 Å. These two scans represent the release of the metal ion to solution and the completion of the second hydration shell, respectively.

The structures with the lowest energy in the beginning and end of the scans were optimized as reactant complex (RC) and product complex (PC), respectively, whereas the structure with the highest energy was optimized as the transition state (TS). After optimization, the dynamic stability of each calculated RC and PC was demonstrated by an absence of negative frequencies; that is, each complex was at least in a local minimum. Possible TSs were identified by the presence of one and only one negative (imaginary) frequency. For each reaction, this negative frequency corresponded either to the forming of the M–OH$_2$ bond or to the breaking of the M–O bond and thus the reaction coordinate between reactants and products. The energy of each RC was set as zero, and the relative energies of the TS and PC were used to make the energy profiles. The PES
scans, geometry optimizations, and frequency calculations were performed with Gaussian 03,\textsuperscript{36} and all images of the reaction mechanisms were made with GaussView 4.\textsuperscript{37}

The H\textsubscript{2}O exchange reaction around the Mg(H\textsubscript{2}O)\textsubscript{6}\textsuperscript{2+} ion was also modeled using the self-consistent reaction field (SCRF) method via the integral equation formalism polarized continuum model (IEFPCM)\textsuperscript{38} with water as the solvent as well as with a super-molecule approach,\textsuperscript{39} where 12 H\textsubscript{2}O molecules were included to model the second hydration sphere. The gas-phase RC, TS, and PC structures were optimized using the IEFPCM. The RC and PC were characterized by the absence of negative frequencies and the TS by a negative frequency corresponding to the formation of the Mg–OH\textsubscript{2} bond. In the super-molecule approach, the starting structure with 6 H\textsubscript{2}O molecules in the first hydration sphere and 12 in the second was optimized. This structure was then used as the input for a PES scan where the distance between the Mg\textsuperscript{2+} and a H\textsubscript{2}O molecule in the second hydration sphere was constrained, and this distance was decreased by 0.1 Å for 20 steps, in a similar fashion to the scans described above. The corresponding lowest energy structures at the beginning and end of the scan were optimized as the RC and PC, respectively, and the highest energy structure was optimized as the TS. As with each of the descriptions given above, the RC and PC structures were characterized by the absence of negative frequencies, while the TS was identified by the presence of one and only one negative frequency corresponding to the formation of the Mg–OH\textsubscript{2} bond.

Once the RC and TS had been isolated for each reaction, the frequency output files were used to calculate the rate constant according to the classical transition state theory (TST) approximation and using TheRate\textsuperscript{40,41} according to the Arrhenius equation (10):
where $A$ is the pre-exponential factor, $T$ is the temperature in K, $n$ is the exponential of the temperature-dependence of $A$, $E_a$ is the activation energy, and $R$ is the gas constant. However, the $n$ exponent was set to zero to eliminate the temperature dependence of $A$, and $T$ was set to 298 K.

### 3.3 Results and Discussion

#### 3.3.1 H$_2$O Exchange around the Hydrated Mg$^{2+}$ Ion

One of the goals of this work is to examine the H$_2$O exchange reaction around metal ions, and the H$_2$O exchange reaction around the Mg(H$_2$O)$_6^{2+}$ ion was studied as a test of our computational approach. Similar investigations have been performed for Ni(H$_2$O)$_6^{2+}$ and Ca(H$_2$O)$_6^{2+}$. To date, computational investigations of hydrated Mg$^{2+}$ ions have been restricted to the stability of structures, hydration energies, binding energies, and vibrational spectra of H$_2$O molecules within the first and second hydration spheres, but no study has focused specifically on determining the barrier height or rate constant of this reaction. Here, H$_2$O exchange around the hexaaqua Mg$^{2+}$ ion is described, and the reaction mechanisms, barrier heights, and rate constants are presented for this reaction in the gas-phase and the aqueous-phase, using the IEFPCM and super-molecule approximations.

The exchange of H$_2$O molecules around the hexaaqua Mg$^{2+}$ ion is shown in Figure 3-1. The energy for each point along the PES scans and the energy profiles for
the gas-phase, IEFPCM, and super-molecule approaches are shown in Figure 3-1a. The reactant complex (RC), transition state (TS), and product complex (PC) structures shown in Figure 3-1b were optimized in the gas-phase. The H$_2$O molecule from the second hydration sphere is circled in blue, while the H$_2$O molecule in the first hydration sphere that will be replaced is circled in green (Figure 3-1b). In the RC, the H$_2$O molecule from the second hydration sphere is H-bonded to two H$_2$O molecules, as has been observed in previous calculations.\textsuperscript{32,46,48} The TS is marked by a single negative frequency corresponding to the formation of the bond between the Mg$^{2+}$ and the H$_2$O molecule in the second hydration sphere. In the PC, the H$_2$O molecule from the second hydration sphere is bonded to the Mg$^{2+}$ ion, and the H$_2$O molecule originally in the first hydration sphere has moved to the second.

The distances between the Mg$^{2+}$ and the approaching second hydration sphere H$_2$O molecule as well as between the Mg$^{2+}$ and the leaving first hydration sphere H$_2$O molecule are listed in Table 3-1 for the optimized RC, TS, and PC structures in the gas-phase. In the RC, the H$_2$O molecule in the second hydration sphere is 3.88 Å from the Mg$^{2+}$, and the H$_2$O molecule in the first hydration sphere is 2.12 Å from the Mg$^{2+}$. As the H$_2$O molecule from the second hydration sphere approaches the Mg$^{2+}$, the bond between the Mg$^{2+}$ and the H$_2$O in the first hydration sphere begins to lengthen. In the TS, this bond length is 2.56 Å, while the H$_2$O from the second hydration sphere is 2.36 Å from the Mg$^{2+}$. Once these two H$_2$O molecules have exchanged, the newly formed Mg–OH$_2$ bond is 2.11 Å, and the H$_2$O that is now in the second hydration sphere is 3.88 Å from the Mg$^{2+}$. 
Figure 3-1: PES scan of H$_2$O exchange around a solvated Mg$^{2+}$ ion. (a) The energy profile for this reaction in the gas-phase and the aqueous-phase. (b) The optimized RC, TS, and PC in the gas-phase. The magnesium ion is yellow, the oxygen atoms are red, and the hydrogen atoms are white. The H$_2$O molecule from the first hydration sphere that moves to the second is circled in green, while the H$_2$O molecule from the second hydration sphere that moves to the first is circled in blue.
Table 3-1: Relevant distances (Å), bond angles (°), barrier heights (kJ/mol), pre-exponential factors (s⁻¹), and rate constants (s⁻¹) for the H₂O exchange reaction around Mg(H₂O)₆²⁺.

<table>
<thead>
<tr>
<th>Optimized Gas-Phase Structure</th>
<th>Mg–H₂O₂nd (Å)</th>
<th>Mg–H₂O₁st (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RC</td>
<td>3.88</td>
<td>2.12</td>
</tr>
<tr>
<td>TS</td>
<td>2.36</td>
<td>2.56</td>
</tr>
<tr>
<td>PC</td>
<td>2.11</td>
<td>3.88</td>
</tr>
</tbody>
</table>

H₂O₂nd–Mg–H₂O₁st Bond Angles in TS (deg)

<p>| | | |</p>
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</thead>
<tbody>
<tr>
<td>Gas-Phase</td>
<td>72, 141, 132, 73</td>
<td></td>
</tr>
<tr>
<td>IEFPCM</td>
<td>91, 110, 92, 107</td>
<td></td>
</tr>
<tr>
<td>Super-Molecule</td>
<td>112, 91, 87, 96</td>
<td></td>
</tr>
</tbody>
</table>

Reaction Phase Barrier Height (kJ/mol) A (s⁻¹) k (s⁻¹)

<p>| | | |</p>
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<tr>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-Phase</td>
<td>71</td>
<td>3.4x10¹³</td>
</tr>
<tr>
<td>IEFPCM</td>
<td>43</td>
<td>5.7x10¹²</td>
</tr>
<tr>
<td>Super-Molecule</td>
<td>33</td>
<td>8.7x10¹¹</td>
</tr>
</tbody>
</table>

The TSs in the gas-phase and aqueous-phase present two different molecular geometries, and they are each pictured in Figure 3-2. The bond angles around Mg²⁺ in the gas-phase, IEFPCM, and super-molecule approximation are given in Table 3-1. For the gas-phase structure, the angles start with the H₂O molecule from the second hydration sphere and proceed clockwise around the Mg²⁺ in Figure 3-2. For the IEFPCM and super-molecule structures, the angles are those made by the axial H₂O molecule, the Mg²⁺ ion, and each equatorial H₂O molecule. In the gas-phase, the molecular geometry is such that the entering and leaving H₂O molecules are oriented 132° from one another, and this contributes to a distorted pentagonal bipyramidal molecular geometry. Despite that this geometry is contrary to expectations,³²,⁴⁶,⁴⁸,⁵⁰-⁵² the heptaqua Mg²⁺ ion has been optimized before.⁴⁸ For both the IEFPCM and super-molecule calculations, the reaction proceeded slightly differently than the gas-phase. Here, the TSs shown in Figure 3-2 have square-pyramidal geometries. In the IEFPCM calculation, the H₂O molecule from the first hydration sphere H-bonded to the H₂O molecule at the apex of the pyramid,
while the \( \text{H}_2\text{O} \) molecule from the second hydration sphere approaches the \( \text{Mg}^{2+} \) along the axis opposite the apex of the pyramid. In the super-molecule calculation, the TS is also square-planar, where the incoming \( \text{H}_2\text{O} \) molecule approaches opposite the axial \( \text{H}_2\text{O} \) molecule bond to the \( \text{Mg}^{2+} \). Due to the presence of the additional, non-reacting \( \text{H}_2\text{O} \) molecules, the first hydration sphere \( \text{H}_2\text{O} \) molecule participates in H-bonding with \( \text{H}_2\text{O} \) molecules in the second hydration sphere rather than those in the first. A square-pyramidal geometry for the TS in \( \text{H}_2\text{O} \) exchange reactions around divalent metal cations has been proposed before.\textsuperscript{33,43,44}

Although the molecular geometries of the TSs are different in the gas-phase and in the aqueous-phase, the \( \text{H}_2\text{O} \) exchange mechanism for \( \text{Mg(H}_2\text{O})_6^{2+} \) most closely matches an associative interchange (I\( _a \)) characterization. The negative frequency in the TS corresponds to the formation of the \( \text{Mg–OH}_2 \) bond and is concerted throughout the H-
bonding network. In the gas-phase and super-molecule calculations, this characterization is further reinforced in that the existing Mg–OH$_2$ bond has not significantly lengthened, < 1.0 Å, in the TS, but this is not true in the IEFPCM calculation. The angles of the octahedron are distorted primarily in the gas-phase calculation. These findings are in contrast to previous work that suggested a seventh group could not be accommodated by Mg$^{2+}$.$^{52,53}$

The PES scans and energy profiles for the Mg(H$_2$O)$_6^{2+}$ exchange reaction in the three phases discussed here are shown in Figure 3-1a, and the barrier heights and rate constants are given in Table 3-1. The plot shows that the gas-phase barrier height is ~30–40 kJ/mol higher than for the IEFPCM and super-molecule calculations, respectively. The barrier heights are 71, 43, and 33 kJ/mol for the gas-phase, IEFPCM, and super-molecule calculations, respectively. The rate constants are calculated according to Equation 3-10 and are 1.1 x 10$^1$, 1.6 x 10$^5$, and 1.7 x 10$^6$ s$^{-1}$, respectively.

The large difference in the gas-phase and aqueous-phase barrier heights likely stems from the hepta-coordinated geometry of the gas-phase TS, and thus the hexa-coordinated geometry of the TSs in the IEFPCM and super-molecule geometries stabilize the reaction. The geometry in the IEFPCM and super-molecule approximation likely results from steric constraints that result from the reaction cavity in the IEFPCM and the additional non-reacting H$_2$O molecules in the super-molecule approach; both of which lower the barrier height. The barrier height of this reaction from the IEFPCM calculation is closest to the experimental values given in Table 3-2, but the super-molecule value is comparable within computational uncertainty. This shows that a molecular cluster
Table 3-2: Reaction enthalpies ($\Delta H^{\ddagger}$, in kJ/mol) and barrier heights ($\Delta E^{\ddagger}$, in kJ/mol) for H$_2$O exchange around Ni(H$_2$O)$_6^{2+}$, Mg(H$_2$O)$_6^{2+}$, and Ca(H$_2$O)$_6^{2+}$ in the gas-phase from previous studies$^{42,43,45,54-56}$ and from this work for Mg(H$_2$O)$_6^{2+}$ in the gas-phase and aqueous-phase.

<table>
<thead>
<tr>
<th></th>
<th>Experimental ($\Delta H^{\ddagger}$, kJ/mol)</th>
<th>Calculated ($\Delta E^{\ddagger}$, kJ/mol)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni$^{2+}$</td>
<td>56.9</td>
<td>42.9</td>
<td>Bechtold, et al., 1978</td>
</tr>
<tr>
<td></td>
<td></td>
<td>46.9</td>
<td>Rotzinger, 1996</td>
</tr>
<tr>
<td></td>
<td></td>
<td>87.4–154.0</td>
<td>Kang, et al., 1991</td>
</tr>
<tr>
<td>Mg$^{2+}$</td>
<td>43</td>
<td>49.1±0.7</td>
<td>Neely and Connick, 1970</td>
</tr>
<tr>
<td></td>
<td>49.1±0.7</td>
<td></td>
<td>Bleuzen, et al., 1997</td>
</tr>
<tr>
<td></td>
<td></td>
<td>71</td>
<td>This work, gas-phase</td>
</tr>
<tr>
<td></td>
<td></td>
<td>43</td>
<td>This work, IEFPCM</td>
</tr>
<tr>
<td></td>
<td></td>
<td>33</td>
<td>This work, super-molecule</td>
</tr>
<tr>
<td>Ca$^{2+}$</td>
<td>28.0, 30.5, 31.4</td>
<td></td>
<td>Tsutsui, et al., 1997</td>
</tr>
</tbody>
</table>

approach to modeling reactions around hydrated metal ions is appropriate for these systems.

The barrier heights for the exchange of H$_2$O molecules around Mg(H$_2$O)$_6^{2+}$ calculated in this work can be compared to the previous experimental and theoretical values given in Table 3-2.$^{42,43,45,54-56}$ The rates of H$_2$O exchange around Ni(H$_2$O)$_6^{2+}$, Mg(H$_2$O)$_6^{2+}$, and Ca(H$_2$O)$_6^{2+}$ are known to follow the series Ni(H$_2$O)$_6^{2+}$ < Mg(H$_2$O)$_6^{2+}$ < Ca(H$_2$O)$_6^{2+}$,$^{57}$ and therefore, one would expect the barrier heights to follow the same trend. Using this logic and the barrier heights in Table 3-2, the barrier heights for H$_2$O exchange around Mg(H$_2$O)$_6^{2+}$ follow the expected trend within computational uncertainty. In addition, the experimentally measured rate constant for this reaction is $6.7 \times 10^5$ s$^{-1}$,$^{57}$ and the aqueous-phase rate constants calculated in this work and given in Table 3-1 are within an order of magnitude of this value.
3.3.2 Reaction Mechanisms and Energy Profiles for H₂O Exchange and Hydrolysis

The PES scans present the energy that each point along the scan has as compared to the energy of the RC, which is set as zero, and the energy profiles represent the energy of the optimized TS and PC species relative to the RC. The scans are presented in Figures 3-3a–7a along with the energy profiles of the optimized RC, TS, and PC, whose structures appear in Figures 3-3b–7b and depict the reaction mechanisms of the five reactions studied. The relevant bond lengths of the optimized RC, TS, and PC as well as the bond angles of the TS for each reaction are given in Table 3-3, and the barrier heights of all the reactions appear in Table 3-4.

3.3.2.1 Protonated Ni–O–Si Sites

The PES scan where the distance between the Ni²⁺ and the H₂O molecule from the second hydration sphere was constrained and served as the reaction coordinate as well as the energy profile appear in Figure 3-3a. The reaction mechanism comprised of the optimized RC, TS, and PC structures is depicted in Figure 3-3b and shows the concerted H₂O exchange and Ni–O bond break. In the RC, the second hydration sphere H₂O molecule is H-bonded to the H⁺ on O as well as to an equatorial H₂O molecule. As this second hydration sphere H₂O molecule approaches Ni²⁺, the equatorial H₂O molecules begin to rearrange to accommodate the second hydration sphere H₂O molecule, as shown in the TS. The negative frequency corresponds to the formation of the Ni–OH₂ bond. As a result, the approach of the H₂O leads to the breaking of the Ni–O bond, and this is
Figure 3-3: (a) The potential energy surface (PES) scan (blue line with blue squares) and optimized energy profile (red line with red squares) for hydrolysis of a protonated Ni–O–Si site where energy (kJ/mol) is plotted versus reaction coordinate. (b) The reaction mechanism for this reaction where the reactant complex (RC), transition state (TS), and product complex (PC) are presented. The nickel ion is blue, the silicon atom is turquoise, the oxygen atoms are red, and the hydrogen atoms are white. The PES scan, geometry optimizations, and frequency calculations were performed by Dr. Amanda A. Olsen during her post-doctoral work at Penn State.
### Table 3-3: Relevant distances (Å) and bond angles (°) in the reactions for protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites.

<table>
<thead>
<tr>
<th>Reaction Type</th>
<th>Optimized Structure</th>
<th>Forming Bond (Å)</th>
<th>Breaking Bond (Å)</th>
<th>Bond Angles (deg)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ni–O–Si</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H₂O Exchange/ Hydrolysis</td>
<td>Ni–H₂O</td>
<td>Ni–O</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RC</td>
<td>3.81</td>
<td>2.10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>2.51</td>
<td>2.14</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC</td>
<td>2.11</td>
<td>4.02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bond Angles (deg)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>O–Ni–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>109, 109, 83, 83</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>H₂O&lt;sub&gt;ax&lt;/sub&gt;–Ni–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>82, 87, 86, 78</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Mg–O–Si</strong></td>
<td>Mg–H₂O&lt;sub&gt;2nd&lt;/sub&gt;</td>
<td>Mg–H₂O&lt;sub&gt;1st&lt;/sub&gt;</td>
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<td></td>
</tr>
<tr>
<td>H₂O Exchange</td>
<td>RC</td>
<td>3.85</td>
<td>2.10</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TS</td>
<td>2.20</td>
<td>2.26</td>
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</tr>
<tr>
<td></td>
<td>PC</td>
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<td>3.84</td>
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<td>Bond Angles (deg)</td>
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<td></td>
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<tr>
<td>TS</td>
<td>O–Mg–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
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<tr>
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<tr>
<td>Hydrolysis</td>
<td>Mg–H₂O</td>
<td>Mg–O</td>
<td></td>
<td></td>
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<tr>
<td>RC</td>
<td>3.85</td>
<td>2.14</td>
<td></td>
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<tr>
<td>TS</td>
<td>4.28</td>
<td>3.74</td>
<td></td>
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<td>PC</td>
<td>2.09</td>
<td>4.11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bond Angles (deg)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>O–Mg–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>70, 105, 72, 70</td>
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</tr>
<tr>
<td>TS</td>
<td>H₂O&lt;sub&gt;ax&lt;/sub&gt;–Mg–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>112, 91, 109, 93</td>
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<td></td>
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<tr>
<td><strong>Ca–O–Si</strong></td>
<td>Ca–H₂O</td>
<td>Ca–O</td>
<td></td>
<td></td>
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<tr>
<td>Hepta-coordinated Ca&lt;sup&gt;2+&lt;/sup&gt;</td>
<td>RC</td>
<td>4.09</td>
<td>2.46</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TS</td>
<td>2.99</td>
<td>2.45</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PC</td>
<td>2.51</td>
<td>2.52</td>
<td></td>
</tr>
<tr>
<td>Bond Angles (deg)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>O–Ca–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>122, 100, 75, 77</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>H₂O&lt;sub&gt;ax&lt;/sub&gt;–Ca–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>84, 93, 84, 77</td>
<td></td>
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</tr>
<tr>
<td>Hydrolysis</td>
<td>Ca–H₂O</td>
<td>Ca–O</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RC</td>
<td>4.09</td>
<td>2.46</td>
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<tr>
<td>TS</td>
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<td>3.04</td>
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<tr>
<td>PC</td>
<td>2.39</td>
<td>4.34</td>
<td></td>
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<tr>
<td>Bond Angles (deg)</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>TS</td>
<td>O–Ca–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>83, 92, 79, 87</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>H₂O&lt;sub&gt;ax&lt;/sub&gt;–Ca–H₂O&lt;sub&gt;eq&lt;/sub&gt;</td>
<td>124, 93, 75, 92</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reaction Type</td>
<td>Barrier Height (kJ/mol)</td>
<td></td>
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<tr>
<td>--------------------------</td>
<td>-------------------------</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Ni–O–Si</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>H₂O Exchange/Hydrolysis</td>
<td>76</td>
<td></td>
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</tr>
<tr>
<td>Mg–O–Si</td>
<td></td>
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<tr>
<td>H₂O Exchange</td>
<td>69</td>
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<tr>
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<tr>
<td>Ca–O–Si</td>
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<tr>
<td>Hepta-Coordinated Ca²⁺</td>
<td>24</td>
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<tr>
<td>Hydrolysis</td>
<td>27</td>
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</tr>
</tbody>
</table>

Table 3-4: Barrier heights (kJ/mol) for H₂O exchange and hydrolysis of protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites.

...evident in the PC by the hexaaqua Ni²⁺ ion and the absence of the Ni–O bond, leaving silicic acid as the second product. The barrier height for this reaction is 76 kJ/mol.

The distances between the Ni²⁺ and the approaching second hydration sphere H₂O molecule and the Ni²⁺ and O are listed in Table 3-3 for the optimized RC, TS, and PC. In the RC, the Ni–H₂O distance is 3.81 Å, and the Ni–O distance is 2.10 Å. For the TS, the Ni–H₂O distance has decreased to 2.51 Å, while the Ni–O distance has increased to 2.14 Å. The final Ni–H₂O and Ni–O distances in the PC are 2.11 Å and 4.02 Å, respectively.

The bond angles surrounding Ni²⁺ in the TS are also included in Table 3-3. The bond angles surrounding the Ni²⁺ in the TS are listed starting from the equatorial H₂O molecule in the right foreground of Figure 3-3b and continue counter-clockwise. The O–Ni–H₂Oeq angles are 109°, 109°, 83°, and 83°, while the H₂Oax–Ni–H₂Oeq bond angles are 82°, 87°, 86°, and 78°. The O–Ni–H₂Oeq angles closest to the approaching second hydration sphere H₂O molecule are increased from the expected 90° for an octahedral...
complex, and this demonstrates that these groups are indeed rearranging as a result of the additional H$_2$O molecule.

The approach of a second hydration sphere H$_2$O molecule toward a protonated Ni–O–Si site leads to release of Ni$^{2+}$, in the form of Ni(H$_2$O)$_6^{2+}$, to solution. The PES scan and mechanism for this reaction indicate that H$_2$O exchange leads to Ni–O bond break. Thus for protonated Ni–O–Si sites, the H$_2$O exchange and hydrolysis reactions are one and the same.

3.3.2.2 Protonated Mg–O–Si Sites

3.3.2.2.1 H$_2$O Exchange

The H$_2$O exchange reaction for a protonated Mg–O–Si site is shown in Figure 3-4. A PES scan of the constrained distance between the Mg$^{2+}$ and the H$_2$O molecule from the second hydration sphere as the reaction coordinate and the energy profile of the optimized RC, TS, and PC appear in Figure 3-4a. The structures of the optimized RC, TS, and PC are shown in Figure 3-4b. As the incoming H$_2$O approaches the Mg$^{2+}$, the equatorial H$_2$O molecules begin to rearrange to enable the exchange. The TS is characterized by elongated bonds to the first hydration shell H$_2$O molecules and to O. Also in the TS, the H$_2$O$_{1st}$ (green circle) begins to leave the first hydration shell, whereas the H$_2$O$_{2nd}$ (blue circle) enters the first hydration shell. For this reaction, the negative frequency corresponds to a concerted interaction of the approaching H$_2$O molecule with the H-bonding network of the equatorial H$_2$O molecules. In the PC, the Mg–O bond is
Figure 3-4: (a) The PES scan (black line with open diamonds) and optimized energy profile (green line with closed diamonds) for H₂O exchange around Mg in a protonated Mg–O–Si site where energy (kJ/mol) is plotted versus reaction coordinate. (b) The reaction mechanism for this reaction where the RC, TS, and PC are presented. The magnesium ion is yellow, the silicon atom is turquoise, the oxygen atoms are red, and the hydrogen atoms are white.
intact, and the H$_2$O$_{1st}$ has been replaced by the H$_2$O$_{2nd}$ as shown in Figure 3-4b. The barrier height for this reaction is 69 kJ/mol.

The distances between the Mg$^{2+}$ and the H$_2$O$_{2nd}$ as well as the Mg$^{2+}$ and H$_2$O$_{1st}$ are listed in Table 3-3 for the optimized RC, TS, and PC in the H$_2$O exchange reaction for the protonated Mg–O–Si site. In the RC, the Mg–H$_2$O$_{2nd}$ distance is 3.85 Å, and the Mg–H$_2$O$_{1st}$ distance is 2.10 Å. In the TS, the Mg–H$_2$O$_{2nd}$ distance is 2.20 Å, shorter than the Mg–H$_2$O$_{1st}$ distance of 2.26 Å. The final Mg–H$_2$O$_{2nd}$ distance is 2.10 Å, and the final Mg–H$_2$O$_{1st}$ distance is 3.84 Å.

The bond angles around Mg$^{2+}$ in the TS are also included in Table 3-3. The bond angles surrounding the Mg$^{2+}$ in the TS are listed using either O or the axial H$_2$O as anchors and starting from the equatorial H$_2$O molecule in the right foreground of Figure 3-4b and continue counter-clockwise. The O–Mg–H$_2$O$_{eq}$ bond angles are 106°, 107°, 82°, and 80°, while the H$_2$O$_{ax}$–Mg–H$_2$O$_{eq}$ angles are 77°, 83°, 89°, and 85°. The O–Mg–H$_2$O$_{eq}$ bond angles show that the equatorial H$_2$O molecules are opening up to accommodate the incoming H$_2$O molecule, and the H$_2$O$_{ax}$–Mg–H$_2$O$_{eq}$ angles show that the equatorial H$_2$O molecules nearest to the approaching H$_2$O molecule are bending toward O also in an effort to accommodate this incoming group.

3.3.2.2.2 Hydrolysis of a Protonated Mg–O–Si Site

Because the stepwise approach of H$_2$O did not lead to the breaking of the Mg–O bond, the possibility of Mg–O bond lengthening as the cause of bond break was examined. The hydrolysis of a protonated Mg–O–Si site appears in Figure 3-5. The
Figure 3-5: (a) The PES scan (blue line with blue diamonds) and optimized energy profile (red line with red diamonds) for hydrolysis of a protonated Mg–O–Si site where energy (kJ/mol) is plotted versus reaction coordinate. (b) The reaction mechanism for this reaction where the RC, TS, and PC are presented. The color scheme is the same as for Figure 3-4.
PES scan is comprised of two steps: the breaking of the Mg–O bond and the absorption of the incoming H₂O molecule by Mg²⁺. The reaction coordinate for the first step is the lengthening of the Mg–O bond, and in the second step, it is the decreasing of Mg–H₂O distance for the second hydration sphere H₂O molecule. The optimized RC, TS, and PC structures comprise the energy profile in Figure 3-5a and are shown as the reaction mechanism in Figure 3-5b. The RC in Figure 3-5b shows the second hydration sphere H₂O molecule H-bonded to the H⁺ on O and an equatorial H₂O molecule, in much the same manner as for the Ni–O–Si site. The Mg–O bond is lengthened until it breaks in the TS, where the negative frequency here corresponds to the breaking of the Mg–O bond, but the second hydration sphere H₂O molecule is not yet absorbed by the Mg²⁺ in the TS. The second hydration sphere H₂O molecule approaches the penta-coordinated Mg²⁺ between the TS and PC and is absorbed in the PC, and thus the PC is comprised of hexaaqua Mg²⁺ ion and silicic acid. The barrier height for this reaction is 54 kJ/mol.

The distances between the Mg²⁺ and the H₂O molecule from the second hydration sphere as well as between the Mg²⁺ and O are listed in Table 3-3. In the RC, the H₂O molecule from the second hydration sphere is 3.85 Å from the Mg²⁺, whereas the Mg–O distance is 2.14 Å. In the TS, the Mg–H₂O distance has increased to 4.28 Å, and the Mg–O distance has increased to 3.74 Å. The increase in the Mg–H₂O distance for the second hydration sphere H₂O molecule increases in the TS in order to allow for the rearrangement of the equatorial H₂O molecules around the Mg²⁺. The final distances for Mg–H₂O and Mg–O are 2.09 Å and 4.11 Å, respectively.

The bond angles around the Mg²⁺ in the TS are also included in Table 3-3. The bond angles surrounding the Mg²⁺ in the TS are listed using either O or the axial H₂O as
anchors and starting from the equatorial H$_2$O molecule in the right foreground of Figure 3-5b and continue counter-clockwise. The O–Mg–H$_2$O$_{eq}$ are 70°, 105°, 72°, and 70°, whereas the H$_2$O$_{ax}$–Mg–H$_2$O$_{eq}$ bond angles are 112°, 91°, 109°, and 93°. These show that the geometry around Mg$^{2+}$ is approaching octahedral, which facilitates the absorption of the H$_2$O molecule from the second hydration sphere.

For protonated Mg–O–Si sites, the H$_2$O exchange and hydrolysis reactions are decoupled. The approach of a H$_2$O molecule causes a H$_2$O from the first hydration sphere to be replaced by one from the second, such that the Mg–O bond break is not a result of the approach of the H$_2$O molecule. Instead, the breaking of the Mg–O bond occurs as a result of the lengthening of the Mg–O bond.

### 3.3.2.3 Protonated Ca–O–Si Sites

#### 3.3.2.3.1 Formation of Hepta-Coordinated Ca$^{2+}$

An examination of H$_2$O exchange for Ca–O–Si sites was performed as well. In this reaction, the second hydration sphere H$_2$O molecule is absorbed by the Ca$^{2+}$ and leads to the formation of a hepta-coordinated Ca$^{2+}$. Thus a true H$_2$O exchange reaction does not occur for this site in these calculations; instead an H$_2$O addition occurs. The PES scan is performed where the incoming H$_2$O molecule approaches the Ca$^{2+}$ (Figure 3-6a). The energies of the optimized RC, TS, and PC are shown in Figure 3-6a and the structures in Figure 3-6b. In the RC, the second hydration sphere H$_2$O molecule is H-bonded to the H$^+$ on O and an equatorial H$_2$O molecule. As the second hydration sphere
Figure 3-6: The (a) PES scan (black line with open triangles) and optimized energy profile (green line with closed triangles) for H₂O exchange around Ca²⁺ in a protonated Ca–O–Si site where energy (kJ/mol) is plotted versus reaction coordinate. (b) The reaction mechanism for this reaction where the RC, TS, and PC are presented. The calcium ion is gold, the silicon atom is turquoise, the oxygen atoms are red, and the hydrogen atoms are white. The PES scan, geometry optimizations, and frequency calculations were performed by Dr. Amanda A. Olsen during her post-doctoral work at Penn State.
H$_2$O molecule approaches the Ca$^{2+}$, the equatorial H$_2$O molecules begin to rearrange to accommodate the incoming H$_2$O molecule as shown in the TS, and the negative frequency corresponds to the formation of this Ca–OH$_2$ bond. However, unlike both the protonated Ni–O–Si and Mg–O–Si sites, the H$_2$O molecule is absorbed to form a hepta-coordinated Ca$^{2+}$, and thus the sole product is this newly-formed calcium silicate cluster. The barrier height for this reaction is 24 kJ/mol.

The Ca–H$_2$O distance for the H$_2$O molecule in the second hydration sphere and the Ca–O distance appear in Table 3-3. In the RC, the Ca–H$_2$O distance is 4.09 Å, and the Ca–O distance is 2.46 Å. The Ca–H$_2$O distance had decreased to 2.99 Å in the TS, while the Ca–O distance has remained nearly constant and is 2.45 Å. In the PC, the Ca–H$_2$O and Ca–O distances are nearly equal at 2.51 Å and 2.52 Å, respectively.

The bond angles surrounding Ca$^{2+}$ in the TS are also included in Table 3-3. The bond angles surrounding the Ca$^{2+}$ in the TS are listed using either the protonated O or the axial H$_2$O as anchors and starting from the equatorial H$_2$O molecule in the right foreground of Figure 3-6b and continue counter-clockwise. The O–Ca–H$_2$O$_{eq}$ bond angles are 122°, 100°, 75°, and 77°, whereas the H$_2$O$_{ax}$–Ca–H$_2$O$_{eq}$ bond angles are 84°, 93°, 84°, and 77°. These angles show that the octahedral geometry around Ca$^{2+}$ is distorting, enabling accommodation of the seventh bonded group.

### 3.3.2.3.2 Hydrolysis of a Protonated Ca–O–Si Site

Because stepwise approach of H$_2$O did not lead to the breaking of the Ca–O bond, the Ca–O bond was lengthened as a possible cause for bond break. The hydrolysis of a
Figure 3-7: (a) The PES scan for the forward direction (blue line with closed triangles), the PES scan for the reverse direction (open blue triangles), and optimized energy profile (red line with closed triangles) for hydrolysis of a protonated Ca–O–Si site where energy (kJ/mol) is plotted versus reaction coordinate. (b) The reaction mechanism for this reaction where the RC, TS, and PC are presented. The color scheme is the same as for Figure 3-6.
protonated Ca–O–Si site appears in Figure 3-7. The initial PES scan appears in Figure 3-7a and is indicated by closed blue triangles and a blue line. The first step in this scan consists of the lengthening and subsequent breaking of the Ca–O bond, whereas the second step is the addition of the H2O molecule to the first hydration sphere. This first attempt to determine the barrier height for the hydrolysis of a protonated Ca–O–Si site led to a barrier height of ~70 kJ/mol, which is higher than the expected value of 25–30 kJ/mol.45

Therefore, the products from this scan were optimized and used to perform a scan of this reaction in the reverse direction, and the reverse direction scan appears as open blue triangles in Figure 3-7a. The TS from this scan was 27 kJ/mol higher in energy than the hexa-coordinated RC from the first scan, making it a representative structure of the overall mechanism. However, the scan in the reverse direction led to the hepta-coordinated Ca-silicate cluster shown in Figure 3-6b. Thus, the energy profile in Figure 3-7a and the reaction mechanism in Figure 3-7b consist of the hexa-coordinated RC, the TS from the reverse scan, and the hexa-coordinated PC.

In the reaction mechanism in Figure 3-7b, the H2O molecule from the second hydration sphere is H-bonded to the H+ on the O and an equatorial H2O molecule. The Ca–O bond is lengthened and eventually breaks in the TS, and the H2O molecule from the second hydration sphere has already been absorbed. The negative frequency in the TS for this reaction corresponds to the breaking of the Ca–O bond. The PC is characterized by an increase in the Ca–O distance, given below, and a fully formed octahedral geometry around Ca2+.  

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The distances between the Ca\(^{2+}\) and the second hydration sphere H\(_2\)O molecule as well as between the Ca\(^{2+}\) and the O appear in Table 3-3. In the RC, the Ca–H\(_2\)O distance is 4.09 Å, and the Ca–O distance is 2.46 Å. The Ca–H\(_2\)O distance decreases to 2.44 Å in the TS, and the Ca–O distance increases to 3.04 Å. The final Ca–H\(_2\)O and Ca–O distances are 2.39 Å and 4.34 Å, respectively, in the PC.

The bond angles around Ca\(^{2+}\) in the TS are also included in Table 3-3. The bond angles surrounding the Ca\(^{2+}\) in the TS are listed using either the H\(_2\)O in the foreground or the H\(_2\)O in the background of Figure 3-7b as anchors and starting from the equatorial H\(_2\)O molecule in the right foreground of Figure 3-7b and continuing counter-clockwise. This method of analysis is used as opposed to the remaining bond angles given in Table 3-3 because here the incoming H\(_2\)O molecule has already been absorbed in the TS. The H\(_2\)O\(_{\text{fore}}\)–Ca–H\(_2\)O\(_{\text{eq}}\) bond angles are 83°, 92°, 79°, and 87°, whereas the H\(_2\)O\(_{\text{back}}\)–Ca–H\(_2\)O\(_{\text{eq}}\) bond angles are 124°, 93°, 75°, and 92°. These angles show that the octahedral geometry around Ca\(^{2+}\) is nearly complete in the TS.

The formation of a hepta-coordinated Ca\(^{2+}\) in the H\(_2\)O exchange reaction for a protonated Ca–O–Si site warrants additional comment. The first hydration shell of Ca\(^{2+}\) has been shown to vary from six to eight.\(^{58-60}\) However, these simulations\(^{58-60}\) and experiments\(^{59}\) were performed for the solution phase and not for the solid state. The molecular clusters used in this work are intended to represent individual sites on a mineral surface, which would be surrounding by many other surface sites. Therefore, the formation of a hepta-coordinated Ca\(^{2+}\) would be unlikely because of steric crowding.

The comparison of H\(_2\)O exchange and hydrolysis for protonated Ca–O–Si sites leads to a third reaction scheme where these two reactions are coupled. The formation of
a hepta-coordinated Ca\(^{2+}\) indicates that the breaking of the Ca–O bond is not affected by
the approach of a H\(_2\)O molecule from the second hydration sphere. This fact is further
reinforced by the elusiveness of the PES scan for the hydrolysis reaction for a protonated
Ca–O–Si site and is manifested by the similarity of the barrier heights and rate constants,
given in Section 3.3.4, for these reactions. The ability of Ca\(^{2+}\) to accommodate a large
number of groups\(^{58,59}\) is what contributes to this complication. However, this elusiveness
likely shows that a specific degree of freedom affects which path is chosen over the other.

3.3.3 Molecular Orbital (MO) and Natural Bond Order (NBO) Analysis of H\(_2\)O
Exchange and Hydrolysis Reactions

The molecular orbitals (MOs) for the hydrolysis reactions are pictured in Figure
3-8, and the RCs and TSs for the reactions at protonated Ni–O–Si, Mg–O–Si, and Ca–O–
Si sites are given. The orbitals surrounding Ni\(^{2+}\) during hydrolysis are comprised of three
bonding orbitals and six anti-bonding orbitals, and the MO pictured in the top row of
Figure 3-8 for Ni–O–Si shows the bonding orbital for the Ni–O bond that will break in
the TS. For protonated Mg–O–Si sites, the MOs surrounding Mg\(^{2+}\) are comprised of one
bonding, one anti-bonding, and three anti-bonding lone pair orbitals. The bonding orbital
pictured in the middle row of Figure 3-8 represents the breaking of the Mg–O bond. For
Ca–O–Si, all of the MOs surrounding Ca\(^{2+}\) in this reaction are anti-bonding. The MO
representing the breaking of the Ca–O bond for the hydrolysis of protonated Ca–O–Si
sites is shown in the bottom row of Figure 3-8, and the MO for Ca–O–Si has extended
electron density on O, the H\(_2\)O in the second hydration sphere, an equatorial H\(_2\)O, and an
axial H\(_2\)O group. For both Ni–O–Si and Mg–O–Si, the breaking of the M–O bond leaves
Figure 3-8: The molecular orbitals (MOs) for the hydrolysis of protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites. The columns correspond to RCs or TSs, and the rows correspond to each silicate site.

an empty space where the second hydration sphere H₂O molecule can bond, and this is evident from the MOs in the TSs for these reactions. For Ca–O–Si, however, the second hydration sphere H₂O molecule has already been absorbed by Ca²⁺, and the electron density in the MO has already rearranged to accommodate the new group. The energies of these orbitals are given in Table 3-5, and they increase in the order Ca²⁺ < Mg²⁺ < Ni²⁺.

The MOs for the H₂O exchange reaction for protonated Mg–O–Si sites and the formation of the hepta-coordinated Ca²⁺ are shown in Figure 3-9. The MOs for the H₂O exchange reaction at a protonated Mg–O–Si site consist of one bonding, one anti-
Figure 3-9: The molecular orbitals (MOs) for the H₂O exchange reaction for protonated Mg–O–Si sites as well as for the formation of the hepta-coordinated Ca–O–Si complex. The columns correspond to RCs or TSs, and the rows correspond to each silicate site.

bonding, and three anti-bonding lone pair orbitals. The MO pictured for this reaction in the top row of Figure 3-9 corresponds to the breaking of the bond between Mg²⁺ and the first hydration sphere H₂O molecule. The MOs for the formation of the hepta-coordinated Ca²⁺ are also pictured in Figure 3-9. For this reaction, the MOs on Ca²⁺ consist of nine anti-bonding orbitals. The MO corresponding to the absorption of H₂O is unlike those in the TSs of the other reactions pictured in Figures 3-8 and 3-9 in that electron density exists on the H₂O molecule from the second hydration sphere and on O. The similar phase on both the Ca²⁺ and the H₂O explains why the H₂O bonds so easily to Ca²⁺, and the breaking of the Mg–H₂O₁st bond in the TS enables the H₂O₂nd to bond with
Table 3-5: Energies (kJ/mol) and change in energy ($\Delta E$, kJ/mol) for relevant molecular orbitals (MOs) in the hydrolysis reactions of protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites, the H$_2$O exchange reaction at the protonated Mg–O–Si site, and the formation of the hepta-coordinated Ca–O–Si site.

<table>
<thead>
<tr>
<th>Protonated Site</th>
<th>RC (kJ/mol)</th>
<th>TS (kJ/mol)</th>
<th>$\Delta E$ (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrolysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ni–O–Si</td>
<td>–286</td>
<td>–288</td>
<td>–2</td>
</tr>
<tr>
<td>Mg–O–Si</td>
<td>–201</td>
<td>–204</td>
<td>–3</td>
</tr>
<tr>
<td>Ca–O–Si</td>
<td>–126</td>
<td>–126</td>
<td>0</td>
</tr>
<tr>
<td>Mg$^{2+}$ H$_2$O Exchange and Hepta-Coordinated Ca$^{2+}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mg–O–Si</td>
<td>–201</td>
<td>–203</td>
<td>–2</td>
</tr>
<tr>
<td>Ca–O–Si</td>
<td>–126</td>
<td>–126</td>
<td>0</td>
</tr>
</tbody>
</table>

Mg$^{2+}$. The energies of these orbitals are presented in Table 3-6, and they show that the energies of the orbitals for Mg$^{2+}$ are higher than those for Ca$^{2+}$.

3.3.4 Rate Constants

The pre-exponential factors and rate constants for the reactions in Reactions 3-4–9 are given in Table 3-6, and the rate constants were calculated using Equation 3-10. The rate constants follow the same trend as the barrier heights outlined above. There are two rate constants for the protonated Mg–O–Si site corresponding to the H$_2$O exchange and hydrolysis reactions, and these rate constants indicate that the hydrolysis reaction would proceed more rapidly than the H$_2$O exchange. Similarly for protonated Ca–O–Si sites, the two rate constants are for the formation of the hepta-coordinated complex and the hydrolysis of this site, where the rate constant of the former is higher than that of the latter. The log of these rate constants are plotted against the log of rate constants for H$_2$O.
Table 3-6: The pre-exponential factors $A \, (s^{-1})$ and rate constants $k \, (s^{-1})$ for H$_2$O exchange and hydrolysis of protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites.

<table>
<thead>
<tr>
<th>Reaction Type</th>
<th>$A , (s^{-1})$</th>
<th>$k , (s^{-1})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni–O–Si</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H$_2$O Exchange/Hydrolysis</td>
<td>$1.7 \times 10^{13}$</td>
<td>$7.2 \times 10^{-1}$</td>
</tr>
<tr>
<td>Mg–O–Si</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H$_2$O Exchange</td>
<td>$3.1 \times 10^{13}$</td>
<td>$2.6 \times 10^{1}$</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>$1.4 \times 10^{14}$</td>
<td>$4.7 \times 10^{4}$</td>
</tr>
<tr>
<td>Ca–O–Si</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepta-Coordinated Ca$^{2+}$</td>
<td>$4.9 \times 10^{14}$</td>
<td>$3.7 \times 10^{9}$</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>$9.3 \times 10^{14}$</td>
<td>$1.5 \times 10^{9}$</td>
</tr>
</tbody>
</table>

exchange around each corresponding metal ion$^{17,57}$ in Figure 3-10, in a similar fashion to the analysis of Casey and Westrich.$^{17}$ The trend is such that the rate constant for the reactions of each metal increases with the rate constant for H$_2$O exchange around the corresponding metal ion.

3.3.5 Overall Trends and Comparison to Experiment

The data in Figure 3-10 replicate the trend observed by Casey and Westrich in experimental data,$^{17}$ where the rates for end-member orthosilicate dissolution increase in the order Ni$^{2+} <$ Mg$^{2+} <$ Ca$^{2+}$. The reaction mechanisms described in this work provide additional insight into the relation of M–O bond energies, H$_2$O exchange rates, and mineral dissolution. The rate constant for the hydrolysis of a protonated surface site increases as the amount of energy required to break the M–O bond decreases, as has been discussed.$^{11,15,16}$ However, what is most intriguing from these calculations is that H$_2$O exchange and hydrolysis reactions are not always intimately related at sites within the clusters, and for some minerals, these phenomena are exclusive for the Mg–O–Si and
Figure 3-10: The log of the rate constant $k$ (s$^{-1}$) for hydrolysis and H$_2$O exchange at protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites (from this work) versus the log of the rate constant $k$ (s$^{-1}$) of H$_2$O exchange for each of the corresponding metal ions.$^{17,57}$

Ca–O–Si linkages. The data in Figure 3-10 also show that the H$_2$O exchange and hydrolysis rate constants are not equal and are likely two reactions occurring within the overall dissolution process.

Although the data in Figure 3-10 replicate the trend of Casey and Westrich,$^{17}$ the values are different. The dissolution rates of Casey and Westrich extend across ~5 orders of magnitude, whereas those in Figure 3-10 differ by ~9 orders of magnitude. There are two likely explanations for this. The higher rate constants calculated here indicate that experimental phenomena contribute to the rate and somehow affect metal release from the surface. Secondly, the experiments of Casey and Westrich are at pH = 2, and although the clusters shown here represent protonated sites on the surface, the
phenomena are different because mineral surfaces contain a distribution of sites in protonated, neutral, and deprotonated states.

The results presented offer a molecular-scale insight into the link between H₂O exchange and the release of metal ions to solution during mineral dissolution. Comparison of the experimental and calculated energies in Table 3-2 for H₂O exchange around Ni(H₂O)₆²⁺, Mg(H₂O)₆²⁺, and Ca(H₂O)₆²⁺ as well as the experimental rate constants given on the x-axis of Figure 3-10 shows that there are differences between the H₂O exchange reactions for these ions and for the reactions described in this work. The presence of a silicic acid group instead of a sixth H₂O molecule around the metal ion is likely responsible for the difference in the H₂O exchange rate constants and the rate constants measured here.⁵¹,⁵²

There are two types of reactions in this regard. The first is where the silicic acid does not participate in the reaction; that is, the M–O bond remains intact. For both the H₂O exchange reaction at the protonated Mg–O–Si site and the formation of the hepta-coordinated Ca²⁺, this silicic acid group affects the rate constant. The H₂O exchange rate constant for Mg(H₂O)₆²⁺ is higher than the rate constant for the reaction in Reaction 3-6, and this shows that the silicic acid kinetically limits the H₂O exchange reaction at a protonated Mg–O–Si site. For the protonated Ca–O–Si site, on the other hand, the rate constant for the formation of the hepta-coordinated Ca²⁺ in the silicate molecular cluster is an order of magnitude larger than that for H₂O exchange on Ca(H₂O)₆²⁺. In this case, the silicic acid group is kinetically enhancing the H₂O exchange.

For the hydrolysis reaction of protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites, the silicic acid group now participates in the reaction; that is, the breaking of the M–O
bond occurs. In these three reactions, the effect of the silicic acid group can also be examined. During hydrolysis, the $\text{H}_2\text{O}$ from the second hydration sphere replaces the silicic acid group, and as a result, the rate constants for these reactions are different than those for $\text{H}_2\text{O}$ exchange reactions for $\text{Ni(H}_2\text{O)}_6^{2+}$, $\text{Mg(H}_2\text{O)}_6^{2+}$, and $\text{Ca(H}_2\text{O)}_6^{2+}$. For both the protonated Ni–O–Si and Mg–O–Si sites, the hydrolysis reactions have smaller rate constants than those for the $\text{H}_2\text{O}$ exchange reactions, whereas for the protonated Ca–O–Si sites, the rate constants for the hydrolysis reaction is larger than that for $\text{H}_2\text{O}$ exchange around $\text{Ca(H}_2\text{O)}_6^{2+}$. These results suggest that the correlation of dissolution for end-member orthosilicate minerals with rate constants of $\text{H}_2\text{O}$ exchange\textsuperscript{17} is coincidental and not an indication that $\text{H}_2\text{O}$ exchange is the mechanism of dissolution.

However, both mineral dissolution and $\text{H}_2\text{O}$ exchange rely on the breaking of M–O bonds and the formation of M–OH$_2$ bonds, and thus the link between M–O bond energy and mineral dissolution rates\textsuperscript{16} is also not surprising.\textsuperscript{11} Furthermore, this connection may show that the bonding environment around the metal ion is similar in the mineral as in the $\text{M(H}_2\text{O)}_6^{2+}$ ion, particularly for transition metals.\textsuperscript{50} Moreover, the occupancy of the $d$ orbitals on Ni$^{2+}$ is the same for both the complexes shown and $\text{Ni(H}_2\text{O)}_6^{2+}$, and the rates of $\text{H}_2\text{O}$ exchange reactions are known to scale with $d$ orbital occupancy for transition metals.\textsuperscript{50,52,53,61} The rate constants calculated also increase with the ionic size of the metal ion, as has been observed in other studies.\textsuperscript{15,17,50-52}

The hydrolysis of protonated Mg–O–Si sites proceeds through a dissociative (D) mechanism. This reaction can be characterized as such because of the decrease in coordination arising from the breaking of the Mg–O bond\textsuperscript{14} and because the Mg–O distance in the TS is > 1.0 Å longer than it was in the RC.\textsuperscript{44} This dissociative type
mechanism is expected for the protonated Mg–O–Si sites based on known mechanisms for H$_2$O exchange around this ion$^{50-52}$ and the primary reason for this expectation is the size of Mg$^{2+}$.\textsuperscript{52,53}

On the other hand, the hydrolysis of protonated Ni–O–Si sites and the H$_2$O exchange reaction at protonated Mg–O–Si sites are in contrast to expectations.\textsuperscript{43,44,50-53} These two reactions appear to proceed through an associative interchange ($I_a$) mechanism for three reasons. First, the negative frequencies correspond to M–H$_2$O bond formation or concerted motion within the H-bonding network. Second, the Ni–O and Mg–H$_2$O$_{1st}$ bonds have not lengthened significantly, < 1.0 Å, in the TS, and third, the bond angles of the octahedron are distorted in the TS. The Ni–O–Si sites are expected to react via dissociative mechanisms because of the population of the $d$ orbitals\textsuperscript{50,52,53,61} and because the approach of the seventh molecule toward the face of the octahedron is electrostatically unfavorable.\textsuperscript{52,53} Also, the small size of the Mg$^{2+}$ ion prevents the incorporation of a seventh group.\textsuperscript{52,53}

However, a recent review of ligand exchange mechanisms showed that Ni$^{2+}$ can react via $D$ or associative ($A$) mechanisms, depending upon the ligands present.\textsuperscript{51} In addition, two previous studies\textsuperscript{43,44} investigated H$_2$O exchange mechanisms around Ni(H$_2$O)$_6^{2+}$ via ab initio calculations. A Hartree-Fock (HF) computational approach was employed because a DFT approach was not possible for transition metals at that time.\textsuperscript{43} A TS for the associative, $A$ or $I_a$, mechanisms,\textsuperscript{43} was not isolated. The TSs presented were square pyramidal in geometry,\textsuperscript{43,44} and the hepta-coordinated TS and intermediate had two negative frequencies.\textsuperscript{43} The author posed the possibility that these results arose as a
results of the method used. These discrepancies in the literature demonstrate that additional investigations are needed.

The mechanism for the formation of the hepta-coordinated Ca\(^{2+}\) can be classified as associative (\(A\)) because the absorption of the incoming H\(_2\)O molecule forms a hepta-coordinated complex without negative frequencies. This is not surprising as an associative mechanism is expected for Ca\(^{2+}\) because of its ionic size.\(^{52}\) One could suggest that perhaps the development of a hepta-coordinated Ca\(^{2+}\) arises because of the employment of a computational approach via molecular clusters. In fact, DFT methods have been shown to prefer decreased coordination numbers for metal ions and thus dissociative mechanisms over associative ones.\(^{61}\) Therefore, the presence of a hepta-coordinated Ca\(^{2+}\) ion in this work is consistent with previous results\(^{58-60}\) and demonstrates this result is not an artifact of gas-phase clusters analyzed by DFT methods.

The hydrolysis of Ca–O–Si sites is difficult to classify. On one hand, the TS is characterized by the breaking of the Ca–O bond, which is indicative of a dissociative interchange (\(I_d\)) mechanism.\(^{51,52,61}\) However, the coordination number of Ca\(^{2+}\) has not decreased. On the other hand, the Ca–OH\(_2\) bond is nearly fully-formed in the TS, and thus an \(A\) mechanism would seem to be an appropriate classification. In this case, the TS would be marked by a negative frequency corresponding to the formation of the Ca–OH\(_2\) bond, which it is not. Thus, the hydrolysis reaction Ca–O–Si sites does not seem to fit into any of the classifications of Langford and Gray,\(^{14}\) and also this reinforces the observation that Ca\(^{2+}\) can react via \(A\) or \(D\) mechanisms.\(^{62}\) Additional analyses are necessary to further elucidate a straightforward description of the PES for the hydrolysis reaction at protonated Ca–O–Si surface sites.
3.4 Conclusions

The H$_2$O exchange reaction around the hexaaqua Mg$^{2+}$ ion is described using gas-phase, IEFPCM, and super-molecule approximations. The barrier height from the IEFPCM calculation matched experimental values most closely, but the value from the super-molecule calculation was comparable within computational uncertainty. The reaction mechanisms are such that the gas-phase TS is characterized by a distorted pentagonal bipyramidal geometry, whereas both the IEFPCM and super-molecule approximations yield square-pyramidal geometries. These differences are manifested in the barrier heights of this reaction, which is ~30–40 kJ/mol higher for the gas-phase. These calculations show that DFT calculations can be used to approximate reactions in aqueous media.

The correlation between H$_2$O exchange reaction rates and the dissolution of end-member silicate minerals has been investigated via DFT methods. Three independent types of reaction mechanisms were determined for protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites. The approach of the H$_2$O molecule from the second hydration sphere leads to the breaking of the Ni–O bond for protonated Ni–O–Si sites, whereas the H$_2$O exchange and hydrolysis reactions are independent for protonated Mg–O–Si sites. For protonated Ca–O–Si sites, however, a third situation develops where the approach of the second hydration sphere H$_2$O molecule is absorbed by the Ca$^{2+}$, forming a hepta-coordinated species. The hydrolysis of protonated Ca–O–Si sites occurs via the lengthening of the Ca–O bond, but PES scans of this reaction were elusive.
The barrier heights from these reactions were used to calculate the rate constants according to the classical TST approximation. The calculated rate constants increase according to \( \text{Ni}^{2+} < \text{Mg}^{2+} < \text{Ca}^{2+} \), which mimics experimental trends.\(^{15,17}\) In addition, the rate constants for the five silicate cluster reactions studied in this work are different than those for \( \text{H}_2\text{O} \) exchange around these ions in solution, and this demonstrates that the presence of a \( \text{Si(OH)}_4 \) group as the sixth ligand instead of a \( \text{H}_2\text{O} \) molecule kinetically affects reactions around these metal ions.\(^{51,52}\) For both protonated \( \text{Ni}–\text{O}–\text{Si} \) and \( \text{Mg}–\text{O}–\text{Si} \) sites, the rate constants for \( \text{H}_2\text{O} \) exchange around \( \text{Ni(H}_2\text{O)}_6^{2+} \) and \( \text{Mg(H}_2\text{O)}_6^{2+} \) are higher than the rate constants for \( \text{H}_2\text{O} \) exchange and hydrolysis. However, for protonated \( \text{Ca}–\text{O}–\text{Si} \) sites, the rate constants for the formation of the hepta-coordinated complex as well as for the hydrolysis reaction are higher than those for \( \text{H}_2\text{O} \) exchange around \( \text{Ca(H}_2\text{O)}_6^{2+} \) in solution.

The reaction coordinates for these reactions were chosen according to whether the approach of the second hydration sphere \( \text{H}_2\text{O} \) molecule led to the breaking of the \( \text{M}–\text{O} \) (\( \text{M} = \text{Ni}, \text{Mg}, \text{or Ca} \)) bond. If not, then the lengthening of the \( \text{M}–\text{O} \) bond was used as the reaction coordinate until it broke, and then a second PES scan was performed to investigate the approach of the second hydration sphere \( \text{H}_2\text{O} \) molecule as the reaction coordinate. Using this approach, the link between \( \text{H}_2\text{O} \) exchange and hydrolysis reactions at protonated sites on end-member silicate minerals was studied.
3.5 References


Chapter 4

Density Functional Theory Modeling of Water on Forsterite (100) and (010) Surfaces

4.1 Introduction

The work builds upon the investigation of Mg$^{2+}$ release from forsterite in Chapter 2. Density functional theory molecular dynamics (DFT-MD) simulations are employed to investigate the structures and relative stabilities of the forsterite (100) and (010) cleavage planes. Instead of studying how the Mg–O bond breaks to effect the release of Mg$^{2+}$ ions to solution, the focus here is to determine the types of functional groups present on the forsterite surface at the aqueous-mineral interface, and surface sites exist as $\text{O}^-$, O$_{\text{br}}$, OH, and H$_2$O groups on both the forsterite (100) and (010) surfaces. Reactions between surface groups and between surface groups and H$_2$O molecules in solution are possible throughout the simulations, and the evolution of surface functional groups over 10 ps is described. Comparisons are given to previous computational and experimental observations.

Characterization of sites on the forsterite surface is necessary for accurate descriptions of dissolution, as the types of functional groups present control the dissolution rate.$^{1,2}$ Currently, the Brunauer-Emmett-Teller (BET) surface area$^3$ is often used to estimate the surface area of a mineral sample, where an inert gas is adsorbed to the mineral surface, and the volume that adsorbs is used to determine the surface area of the mineral. This term is then used to estimate the surface contribution to the dissolution
rate law. The overall dissolution rate does not correlate well with the BET surface area in many instances, and as a result, chemical probe molecules are being used to determine which sites are the most reactive on mineral surfaces. In addition, the adsorption of the inert gas during a BET surface area measurement is not sensitive to surface topographical features, and this serves as another disadvantage to the technique in that surface topography also contributes to the dissolution rate of a mineral because preferential dissolution occurs at defect sites and edge sites.

In addition to descriptions of which functional groups are present on the forsterite mineral surface, the relative stabilities of crystallographic planes also contribute to observed dissolution rates. The two major reasons for this are that each crystallographic plane will be present in varying concentrations and because each plane will be terminated with a different number of each functional group. The relative stabilities of forsterite (100) and (010) surfaces have been investigated using atomistic simulations. The means by which the forsterite surface is cleaved affects which surface is determined to be more stable, as too many dangling bonds in the cleavage plane can lead to a surface that is heavily covered in defects. This difference in cleavage of the surface plane may be why the (100) surface was found to be more stable in one study and less stable in another. These conflicting conclusions demonstrate that additional insight is needed to isolate which of these two surfaces is more stable than the other, and such a conflict can be resolved with the use of density functional theory molecular dynamics (DFT-MD) simulations.

Adsorption of water onto the forsterite mineral surface has several overarching applications. Forsterite is one of the most common astronomical minerals, and
therefore, understanding the structure of monolayers of water on the forsterite mineral surface provides insight into a number of astronomical phenomena. The first is how water was delivered to Earth,\textsuperscript{12,14,15} as water is believed to have adsorbed onto forsterite grains during planet formation.\textsuperscript{15} In addition, the adsorption of water onto forsterite influences the current concentration of water in Earth’s mantle,\textsuperscript{16-20} and the presence of water affects seismic processes\textsuperscript{17-19} and the global water cycle\textsuperscript{18} throughout Earth’s surface. Lastly, forsterite has been found on the surface of Mars, and thus, understanding the structure of water on the surface of this mineral enables a description of the mineralogical history of this planet.\textsuperscript{21-23}

DFT-MD simulations for the investigation of the surface structure of the forsterite (100) and (010) planes in both hydrated and aqueous environments are particularly useful because both bond-breaking and bulk scale behaviors can be studied simultaneously. DFT-MD simulations via the Vienna Ab-initio Simulation Package (VASP)\textsuperscript{24-27} have been employed to model the adsorption of H\textsubscript{2}O molecules to mineral surfaces,\textsuperscript{28} the aqueous-mineral interface,\textsuperscript{29,30} the adsorption of ions from solution to the mineral surface,\textsuperscript{31} and the vibrational spectrum of H\textsubscript{2}O molecules adsorbed to the mineral surface.\textsuperscript{32} The advantage of using DFT-MD versus classical MD in part is that the results from the simulations are not artifacts of the potentials or fitting parameters used.\textsuperscript{29} In addition, reactions between surface species and solution species can occur in DFT-MD, whereas many classical force fields rely on non-dissociable H\textsubscript{2}O molecules. Consequently, DFT-MD allows for a more realistic description of the surface and is less subject to the initial configuration selected. This approach builds upon previous work of
using single surface sites to describe the release of Mg$^{2+}$ from the forsterite surface by increasing the number of surface sites and modeling cooperative H-bonding effects.

The goal of this work is to determine the structure of the forsterite (100) and (010) surfaces in both H$_2$O monolayer and bulk aqueous environments. The energetic stability of each will be calculated via DFT-MD simulations with the intent that these simulations will provide insight for experimental characterization and dissolution studies. Use of quantum mechanical calculations enables bond-breaking and bond-forming processes to be included in the simulations, and thus H$^+$ and H$_2$O transfers between surface groups and at the aqueous-mineral interface are possible. Each surface is investigated at a unit cell scale which permits the inclusion of several surface sites, and thus the protonation states on the surfaces in hydrated and aqueous environments will be determined. The findings of this study are then compared with previous computational and experimental results.

4.2 Computational Details

Density functional theory molecular dynamics (DFT-MD) simulations have the benefits of including the electronic structure of atoms as well as long enough timescales for the investigation of physicochemical processes. Simulations are made practical by employing pseudopotentials that separate the non-reacting core electrons from the valence electrons during the calculation of the electronic structure. The Vienna Ab-initio Simulation Package (VASP) employs a plane wave approximation to the band structure via the projector augmented-wave (PAW) approach. The advantages of the
PAW approach are the elimination of adjustments to the core electrons and the ability to achieve chemical behaviors similar to those observed in all-electron methods. The improvements by Kresse and Joubert on Blöchl’s original description are especially useful for silicate systems because the properties of a siliceous substance were tested in the training set. The calculations described in this work were constructed with an automatic generation of \( k \) points within the Brillouin zone according to the Monkhorst-Pack approach. The energy cutoff for the energy-minimization of samples was 500 eV, which is a reasonably accurate cut-off for these systems.

An overview of the computational scheme appears in Figure 4-1. The coordinates for the forsterite (100) and (010) surfaces were taken from the Cerius² mineral structures database, and the bulk crystal was energy-minimized such that the periodic cell boundaries were permitted to change. The force and stress tensors were calculated. Simultaneously, the positions of all ions were permitted to relax. The coordinates of the bulk model were used as the lattice parameters for further surface simulations, and the energy of this energy-minimized structure is \( E_{bulk} \).

Surface slabs were cleaved from the forsterite mineral structure using the Surface Builder module in the Cerius² database along the (100) and (010) cleavage planes in such a way as to minimize the number of dangling bonds. The forsterite (100) slab is 10.2430 x 6.0030 Å² and is 11.75 Å thick, and this cell has eight Mg₂SiO₄ formula units. The forsterite (010) slab is 12.0060 x 9.5380 Å² and is 15.75 Å thick, and this cell has 16 formula units. These slabs were also energy-minimized allowing all ions to relax but restricting the dimensions of the simulation cell. The final dimensions of the forsterite (100) cell were 19.6954 x 10.2430 x 6.0030 = 1211.0 Å³, and the forsterite (010) cell was
21.3124 x 12.0060 x 9.5380 = 2440.6 Å³. The energy of these energy-minimized slabs are denoted $E_{slab}$.

Each surface was then covered with H$_2$O or H$^+$ and OH$^-$ groups to model associative and dissociative adsorption, respectively. A neutral charge in the system was maintained by adjusting the number of H$^+$ and OH$^-$ groups bonded to the surface. These model surfaces were energy-minimized, and again all ions were permitted to relax. The free energy of the energy-minimized monolayer samples is $E_{ads}$. The H$_2$O molecules adsorbed to each surface in the monolayer system were energy-minimized separately in a cell whose dimensions were the same as the vacuum portion of the forsterite (100) and
(010) simulation cells, and the free energy of this energy-minimized configuration is labeled as $E_{\text{hyd}}$.

The surface energy indicates the relative stability of a surface structure, and the values calculated in this work will be used to determine whether the forsterite (100) or (010) surface is more stable for the slabs, hydrated surfaces, and at the aqueous-mineral interface. According to the description of de Leeuw et al., the surface energy $\gamma$ of a mineral slab is calculated according to (1):

$$\gamma = \frac{E_{\text{slab}} - E_{\text{bulk}}}{A}$$  \hspace{1cm} (1)

where $A$ is the area of the surface (in units of m$^2$). For all the samples in this work, $A$ is multiplied by a factor of two to account for both sides of the slab, as both are permitted to react as a result of the periodic boundary conditions. $E_{\text{slab}}$ and $E_{\text{bulk}}$ are each reflective of the number of Mg$_2$SiO$_4$ formula units within the slab and bulk, respectively. The relative energy of the hydrated slabs are calculated according to (2).

$$\gamma = \frac{E_{\text{ads}} - (E_{\text{bulk}} + E_{\text{hyd}})}{A}$$  \hspace{1cm} (2)

An additional system was designed to model the low density of water that could exist if vapor-phase H$_2$O adsorbed onto the surface. In this system, the energy-minimized (100) surface covered with a distribution of functional groups, formed from six adsorbed H$_2$O molecules, was solvated by 12 H$_2$O molecules to simulate the aqueous-mineral interface. The six H$_2$O molecules used to hydrate the surface and the 12 H$_2$O molecules used to solvate the surface were separately energy-minimized in cells that were the same size as the vacuum above the (100) surface. The energies of these molecules
are as listed as $E_{\text{hyd}}$ and $E_{\text{solv}}$ in Table 4-1, respectively. DFT-MD simulations were performed with a 0.5 fs timestep for a total of 10 ps, and the final 5 ps were used for data analysis. The advantage to this model is that less H$_2$O molecules within the simulation cell allow for more freedom of H$_2$O arrangement. Following similar logic to de Leeuw et al., the relative energy of this low density system was calculated according to (3):

$$\gamma = \frac{E_{\text{int}} - (E_{\text{bulk}} + E_{\text{hyd}} + E_{\text{ads}} + E_{\text{solv}})}{A}$$

The models studied in this work were visualized with either Cerius$^2$ or Materials Studio.$^{39}$

4.3 Results and Discussion

4.3.1 Bulk Forsterite Structure

The original and energy-minimized structures for bulk forsterite are shown in Figure 4-2 and correspond to 16 formula units. The lattice parameters of the original sample were (9.538 x 10.243 x 12.006) Å$^3$, and those for the DFT energy-minimized structures were (9.600 x 10.321 x 12.095) Å$^3$. The errors in calculated lattice parameters of > 1% are excellent for the DFT methodology used here.$^{40}$ The angles of the energy-minimized bulk structure are $\alpha = \beta = \gamma = 90^\circ$. The energy-minimized structure has a free energy of $-4.65 \times 10^3$ kJ/mol, and this value is listed as $E_{\text{bulk}}$ in Table 4-1.
Table 4-1: Energies (kJ/mol) and lattice parameters (Å³) for forsterite samples studied throughout this work.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Lattice Parameters</th>
<th>Energy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bulk</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Input</td>
<td>9.538 x 10.243 x 12.006</td>
<td></td>
</tr>
<tr>
<td>Output</td>
<td>9.600 x 10.321 x 12.095</td>
<td>−4.65x10³</td>
</tr>
<tr>
<td><strong>Hydrated Forsterite (100)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Slab</td>
<td>−4.48x10³</td>
<td></td>
</tr>
<tr>
<td>Associative Adsorption</td>
<td>8 H₂O molecules</td>
<td>−1.13x10⁴</td>
</tr>
<tr>
<td></td>
<td>Slab + Adsorbed H₂O</td>
<td>−8.2x10⁴</td>
</tr>
<tr>
<td>Dissociative Adsorption</td>
<td>8 H₂O molecules</td>
<td>−1.13x10⁴</td>
</tr>
<tr>
<td></td>
<td>Slab + Adsorbed H₂O</td>
<td>−8.31x10⁴</td>
</tr>
<tr>
<td><strong>Low-Density Solvated Forsterite (100)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aqueous-Mineral Interface</td>
<td>6 H₂O molecules</td>
<td>−8.43x10³</td>
</tr>
<tr>
<td></td>
<td>Slab + Adsorbed H₂O</td>
<td>−4.54x10⁴</td>
</tr>
<tr>
<td></td>
<td>12 H₂O molecules</td>
<td>−1.70x10⁴</td>
</tr>
<tr>
<td></td>
<td>Hydrated slab +</td>
<td>−6.22x10⁴</td>
</tr>
<tr>
<td></td>
<td>12 H₂O molecules</td>
<td></td>
</tr>
<tr>
<td><strong>Hydrated Forsterite (010)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Slab</td>
<td>−4.39x10³</td>
<td></td>
</tr>
<tr>
<td>Associative Adsorption</td>
<td>12 H₂O molecules</td>
<td>−1.70x10⁴</td>
</tr>
<tr>
<td></td>
<td>Slab + Adsorbed H₂O</td>
<td>−8.81x10⁴</td>
</tr>
<tr>
<td>Dissociative Adsorption</td>
<td>8 H₂O molecules</td>
<td>−1.13x10⁴</td>
</tr>
<tr>
<td></td>
<td>Slab + Adsorbed H₂O</td>
<td>−8.31x10⁴</td>
</tr>
</tbody>
</table>
4.3.2 Forsterite (100) and (010) Slabs

Surface structures are shown in Figures 4-3 and 4-4 for the forsterite (100) and (010) cleavage planes, respectively. The numbers of functional groups on the forsterite (100) and (010) slabs are given in Table 4-2. The initial (100) cleavage plane contained five O br and two O − functional groups on the surface, where the total number of sites possible was 36. Upon energy-minimization, all of the O atoms on the surface became O br groups. The initial (010) sample, on the other hand, had 28 O br and four O − out of a possible 80 sites on the surface. The energy-minimized (010) cleavage plane had 28 O br and four O − functional groups, which is the same as the original sample. The total possible sites refers to the number of dangling surface bonds on both Mg $^{2+}$ ions and Si.
Figure 4-3: Forsterite (100) slab. (a) Initial structure.\textsuperscript{37,39} (b) Energy-minimized structure.\textsuperscript{39} The color scheme is the same as Figure 4-1.

Figure 4-4: Forsterite (010) slab. (a) Initial structure.\textsuperscript{37,39} (b) Energy-minimized structure.\textsuperscript{39} The color scheme is the same as Figure 4-1.
atoms on the surface of these mineral samples, and this terminology will be continued throughout this work.

The energies of the energy-minimized slabs are listed in Table 4-1 as $E_{\text{slab}}$. The total energy for the forsterite (100) slab was $-4.48 \times 10^3$ kJ/mol, and the total energy for the forsterite (010) slab was $-4.39 \times 10^3$ kJ/mol. Each slab had 8 and 16 Mg$_2$SiO$_4$ formula units for the (100) and (010) slabs, respectively.

The initial structure of the (100) cleavage plane was such that atoms were more free to move, and thus, the surface underwent significant rearrangement during energy-minimization. This observation is in contrast to what was seen for the (010) cleavage plane, where the configuration of atoms on the surface did not enable significant rearrangement. This may in part explain why a lower energy structure was attained for the (100) cleavage plane as a result.

The relative energies $\gamma$ of the (100) and (010) slabs are given in Table 4-3. For the (100) slab, $\gamma$ is 1.81 J/m$^2$, while for the (010) slab, $\gamma$ is 3.07 J/m$^2$. Thus, the (100) surface is predicted to be more stable than the (010) surface in vacuum because of the lower surface energy.

### 4.3.3 Associative and Dissociative Adsorption of H$_2$O to Forsterite Slabs

The adsorption of H$_2$O molecules to the forsterite (100) and (010) surfaces was modeled to determine whether associative or dissociative adsorption was more favorable for each plane. The energy values for these samples are given in Table 4-1 as $E_{\text{ads}}$, and the number of functional groups on each respective original and energy-minimized
Table 4-2: Type and number of functional groups on the bulk, slab, hydrated, and aqueous-mineral interface surfaces for forsterite (100) and (010). The raw number and the fraction of total sites are given.

<table>
<thead>
<tr>
<th>Adsorption</th>
<th>Structure</th>
<th>O_{br}</th>
<th>O^{-}</th>
<th>OH</th>
<th>H_{2}O</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slab</td>
<td>Hydrated Forsterite (100): 36 total possible surface sites</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Original</td>
<td>5 (14%)</td>
<td>2 (6%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td></td>
<td>Energy-minimized</td>
<td>7 (19%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Associated</td>
<td>Original</td>
<td>6 (17%)</td>
<td>2 (6%)</td>
<td>0 (0%)</td>
<td>8 (22%)</td>
</tr>
<tr>
<td></td>
<td>Energy-minimized</td>
<td>3 (8%)</td>
<td>0 (0%)</td>
<td>10 (28%)</td>
<td>2 (6%)</td>
</tr>
<tr>
<td>Dissociated</td>
<td>Original</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>16 (44%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td></td>
<td>Energy-minimized</td>
<td>2 (6%)</td>
<td>0 (0%)</td>
<td>12 (33%)</td>
<td>2 (6%)</td>
</tr>
<tr>
<td>Low-Density Solvated Forsterite (100): 36 total possible surface sites</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aqueous Interface</td>
<td>t = 0 ps</td>
<td>2 (6%)</td>
<td>1 (3%)</td>
<td>10 (28%)</td>
<td>2 (6%)</td>
</tr>
<tr>
<td></td>
<td>t = 10 ps</td>
<td>2 (6%)</td>
<td>2 (6%)</td>
<td>9 (25%)</td>
<td>3 (8%)</td>
</tr>
<tr>
<td>Hydrated Forsterite (010): 80 total possible surface sites</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Slab</td>
<td>Original</td>
<td>28 (35%)</td>
<td>4 (5%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td></td>
<td>Energy-minimized</td>
<td>28 (35%)</td>
<td>4 (5%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Associated</td>
<td>Original</td>
<td>28 (35%)</td>
<td>4 (5%)</td>
<td>0 (0%)</td>
<td>12 (15%)</td>
</tr>
<tr>
<td></td>
<td>Energy-minimized</td>
<td>26 (33%)</td>
<td>4 (5%)</td>
<td>4 (5%)</td>
<td>8 (10%)</td>
</tr>
<tr>
<td>Dissociated</td>
<td>Original</td>
<td>24 (30%)</td>
<td>0 (0%)</td>
<td>16 (20%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td></td>
<td>Energy-minimized</td>
<td>25 (31%)</td>
<td>0 (0%)</td>
<td>14 (18%)</td>
<td>1 (1%)</td>
</tr>
</tbody>
</table>

Table 4-3: Relative energies (J/m²) for the forsterite (100) and (010) slabs, hydrated surfaces, and aqueous-mineral interfaces.

<table>
<thead>
<tr>
<th>Surface Plane</th>
<th>Slab</th>
<th>Hydrated Surface</th>
<th>Aqueous-Mineral Interface</th>
</tr>
</thead>
<tbody>
<tr>
<td>(100)</td>
<td>Pore Water System</td>
<td>1.8</td>
<td>0.32</td>
</tr>
<tr>
<td>(010)</td>
<td>3.1</td>
<td>2.47</td>
<td>1.89</td>
</tr>
</tbody>
</table>
Figure 4-5: Hydrated forsterite (100) surface. a) Original (left) and energy-minimized (right) surface structures for associative adsorption of H₂O molecules. b) Original (left) and energy-minimized (right) surface structures for dissociative adsorption of H₂O molecules. The color scheme is such that Mg atoms are yellow, Si atoms are green, O atoms are red, and H atoms are white.\textsuperscript{37,39}
surface is listed in Table 4-2. For the (100) plane, associative adsorption and dissociative adsorption of H₂O molecules were modeled with eight H₂O molecules adsorbed to the slab where the total energy contribution from the H₂O molecules adsorbed (\(E_{\text{hyd}}\)) was –1.12 x 10⁴ kJ/mol. The initial and energy-minimized surfaces are shown in Figure 4-5. The energy- minimization of the associative adsorption surface was –4.82 x 10⁴ kJ/mol, and during the energy-minimization for associative adsorption, one H₂O molecule was released from the surface to the vacuum. For dissociative adsorption, on the other hand, the energy-minimization of the dissociative adsorption surface was –4.83 x 10⁴ kJ/mol. Although these two values are nearly identical, dissociative adsorption is predicted to be a lower energy process for the forsterite (100) surface because a surface H₂O molecule was released during the energy-minimization of associative adsorption.

The associative and dissociative mechanisms of H₂O adsorption to the forsterite (010) surface were also modeled, and these surfaces are shown in Figure 4-6. For the (010) plane, 12 intact H₂O molecules were adsorbed to the slab, where the total energy contribution from the H₂O molecules adsorbed was –1.70 x 10⁴ kJ/mol. The energy- minimization of the associative adsorption surface was –8.81 x 10⁴ kJ/mol, and during this simulation, two H₂O molecules were released from the surface to the vacuum. For dissociative adsorption, on the other hand, eight H₂O molecules were adsorbed to the forsterite (010) slab, and the energy contribution of these molecules was –1.13 x 10⁴ kJ/mol. Similar to the associative adsorption for this cleavage plane, one H₂O molecule was released from the surface to the vacuum. Two different numbers of H₂O molecules were adsorbed for the (010) cleavage plane to maintain charge neutrality in the system,
Figure 4-6: Hydrated forsterite (010) surface. a) Original (left) and energy-minimized (right) surface structures for associative adsorption of H₂O molecules. b) Original (left) and energy-minimized (right) surface structures for dissociative adsorption of H₂O molecules. The color scheme is the same as Figure 4-5. 37,39
which was a challenge due to the structure of the plane, particularly in the dissociative adsorption case. The energy-minimization of the dissociative adsorption surface was \(-8.31 \times 10^4\) kJ/mol. Because the cells used to simulate dissociative and associative adsorption have different numbers of atoms, comparisons regarding the energy of these processes will be limited to the relative energies, discussed below.

The relative energies for the associative and dissociative adsorption of \(\text{H}_2\text{O}\) molecules on the (100) and (010) cleavage planes are given in Table 4-3. For the (100) cleavage plane, \(\gamma_{\text{diss}}\) is 0.26 J/m\(^2\), and \(\gamma_{\text{assoc}}\) is 0.32 J/m\(^2\). As before, the similarity of these two values is a result of the nearly identical energy values for the hydrated surfaces. For the (010) cleavage plane, \(\gamma_{\text{diss}}\) is 1.89 J/m\(^2\), and \(\gamma_{\text{assoc}}\) is 2.47 J/m\(^2\). This confirms the expectation that dissociative adsorption would be a lower energy process for this cleavage plane.

4.3.4 Density Functional Theory Molecular Dynamics (DFT-MD) Simulations

The results outlined in Sections 4.3.1–4.3.3 serve as the foundation for the simulations described in the current section. In the above analysis, the goals were to determine whether the (100) cleavage plane was more stable than the (010) cleavage plane and to investigate the surface structure of these planes when a monolayer of \(\text{H}_2\text{O}\) molecules is adsorbed to them. These systems were used to make samples that represented the aqueous-mineral interface for the forsterite (100) and (010) surfaces, where MD simulations were performed to investigate how the distribution of functional
groups on the surface changes with time and how the H-bonding network at the aqueous-mineral interface affects the surface structure.

The hydrated forsterite (100) surface that is solvated by 12 H$_2$O molecules is characterized by one O$^-$, two H$_2$O, ten OH, and two O$_{br}$ groups as listed in Table 4-2, and the initial starting configuration for the aqueous-mineral interface of this plane is shown in Figure 4-7a. There are 110 atoms in the simulation cell. The x-coordinates correspond to the vertical direction of Figure 4-7a, and the assignments of regions as surface, bulk water, and slab for forsterite are also shown. The average positions of O and H atoms in the forsterite (100) simulation cell for the low-density water system are shown in Figures 4-7b and 4-7c, respectively. As with the fully-solvated forsterite (100) system, the locations of the O atoms remain roughly constant, with the exception of the O atoms in H$_2$O molecules in solution. For the H atom histogram shown in Figure 4-7c, the wide peaks from $0.00 \text{ Å} \leq x \leq 10.25 \text{ Å}$ show that the H atoms contained within this part of the system move quite a bit during the last 5 ps of the simulation.

The data described above for the forsterite (100) solvated by 12 H$_2$O molecules average the positions of O and H atoms over time. Throughout the simulation, the O$^-$ group on the surface remained deprotonated except for a H$^+$ transfer from a neighboring H$_2$O on the surface, but this H$^+$ is transferred back within 0.5 ps. In addition, a H$^+$ from an OH group on the surface is transferred to another, leaving O$^-$ and H$_2$O groups on the surface at the end of the simulation. The O$_{br}$ remains deprotonated throughout the entirety of the simulation, however. After 10 ps, there are three H$_2$O molecules adsorbed to the forsterite (100) surface for the low-density water system, and this corresponds to a
Figure 4-7: (a) Forsterite (100) surface solvated with low-density water. Atoms are characterized according to their $x$-coordinates (b) O atom histogram and (c) H atom histogram, where the absence of H atoms from $11 \leq x \leq 20$ corresponds to the forsterite slab. Each histogram is averaged over the last 5 ps of simulation. The color scheme is the same as Figure 4-5.
coverage of ~2 H₂O/nm². The relative stability of the forsterite (100) aqueous-mineral interface with 12 H₂O molecules studied here is 61.8 J/m² (Table 4-3).

4.3.5 Comparison to Computational Results and Experimental Data

The bulk crystal lattice dimensions calculated in this work are within 1% of the experimentally measured lattice parameters of \( a = 4.756 \, \text{Å}, \ b = 10.207 \, \text{Å}, \ c = 5.980 \, \text{Å} \).\(^{41}\) In addition, the forsterite unit cell angles were determined to be \( \alpha = \beta = \gamma = 90^\circ \) for a forsterite sample used in recent experiments,\(^{42}\) and the angles for the energy-minimized bulk sample from this work were the same as the experimental sample. The comparison of lattice parameters shows that the VASP calculations described in this work model the
electronic structure of this system well. Therefore, our computational methodology is an appropriate method to study forsterite surfaces.

The relative stabilities of the forsterite (100) and (010) surfaces were also calculated and are included in Table 4-3. The forsterite (100) surface was found to be more stable than the (010) surface, and this agrees with earlier findings.\(^9\) In addition, the dissociative adsorption surfaces were slightly more stable than the corresponding associative adsorption surfaces for the same cleavage plane. This is likely because less energy was required to stabilize the surfaces that already had H\(_2\)O molecules in a dissociated state. Furthermore, both the associative and dissociative adsorption samples minimize to surfaces with a similar distribution of functional groups on the surface.

These findings contrast earlier investigations employing force field calculations\(^{38}\) and DFT calculations\(^{43}\) where the authors concluded that associative and dissociative adsorption happen exclusively. Further, associative adsorption was considered more favorable because the process was calculated to be more exothermic.\(^{38}\) However, this was a result of the fact that the H\(_2\)O molecules on the forsterite surface were not able to dissociate or recombine in those calculations because a non-reactive force field was used. At the same time, the H\(_2\)O molecules were found to prefer Mg\(^{2+}\) sites in the samples simulated here, which served to complete the coordination sphere on the forsterite surface, and similar results were found in a previous study.\(^{38}\) More recently, density functional theory (DFT) calculations showed that dissociated H\(_2\)O molecules recombine,\(^{43}\) which is also in accord with our current findings.

For both the forsterite (100) and (010) cleavage planes, the energy-minimized forms of the surfaces initially covered with either intact or dissociated H\(_2\)O molecules are
similar. A distribution of O\textsuperscript{-}, O\textsubscript{br}, OH, and H\textsubscript{2}O groups exists on each. These observations show that the protonation state of sites is an important contribution for the forsterite mineral structure. In addition, not all sites on the forsterite surface are covered with OH groups, which provides information regarding where dissolution reactions occur on the surface of this mineral\textsuperscript{2,44} The results described show that different types of sites exist on the forsterite mineral surface, which corroborates the idea of identifying reactive sites on a mineral surface versus the total number of sites and total surface area\textsuperscript{4,45}.

A recent description of Mg\textsuperscript{2+} release from the forsterite mineral surface used the total number of OH groups possible on the (100) plane as possible reactive sites\textsuperscript{2}. The rate of Mg\textsuperscript{2+} release calculated was \(\sim 10^6\)–\(10^9\) times faster than experimentally measured rates, and the authors attributed this discrepancy to the possibility that they were not modeling the rate-limiting step\textsuperscript{2}. However, another possibility is that the total number of OH groups does not equal the number of sites where a dissolution reaction will occur. As seen in this surface characterization study, not every functional group on the forsterite (100) and (010) surfaces is an OH group. What is more, OH sites are known to be preferential dissolution sites\textsuperscript{46}.

If one combines the results of this work with those of Morrow et al\textsuperscript{2}, then a new dissolution rate can be calculated. Using either the actual number of OH groups on the hydrated forsterite (100) or (010) surfaces or the number of H\textsubscript{2}O and O\textsuperscript{-} groups, the Mg\textsuperscript{2+} release rate decreases by less than an order of magnitude (data not shown). This suggests that the rate of Mg\textsuperscript{2+} release is not only controlled by the types of groups on the surface but more importantly by the reactions that ultimately lead to surface functionality, such as degradation of the mineral structure as a result of adsorption of groups to the surface\textsuperscript{47}.
In addition, the formation of a silica rich layer on the surface of forsterite during dissolution\textsuperscript{42,48} also contributes to the overall rate for \( \text{Mg}^{2+} \) release.\textsuperscript{2}

4.4 Conclusions and Future Directions

4.4.1 Conclusions

The Vienna Ab-initio Simulation Package (VASP) was used to model the adsorption of \( \text{H}_2\text{O} \) molecules onto the forsterite (100) and (010) cleavage planes. For both dissociative and associative adsorption, a distribution of sites that includes \( \text{O}^- \), \( \text{O}_{br} \), \( \text{OH} \), and \( \text{H}_2\text{O} \) groups results for the energy-minimized surfaces. However, dissociative adsorption is predicted to be a lower energy process because the resulting surface was lower in energy than that for the associative surface for both the forsterite (100) and (010) cleavage planes. These results provide insight into reactive surface sites for this mineral in that not all of the sites on the surface are \( \text{OH} \) groups, as was previously modeled.\textsuperscript{2}

The forsterite (100) and (010) cleavage planes were compared to identify which was more stable, and forsterite (100) was determined to be the more stable plane. This is in accord with previous force field calculations\textsuperscript{9} but in contrast to another.\textsuperscript{38} In the present work, particular attention was given to cutting the slabs so that the number of dangling bonds was minimized, which can lead to the development of surface defects.\textsuperscript{38} Knowledge of which plane is more stable is important for determining the concentration of each plane in a mineral sample, as the more stable planes are present in higher concentration.
The interaction of water with the forsterite surface as a monolayer phase has implications for the incorporation of water into the Earth during its accretionary phase. Currently, the means by which water arrived on Earth is not understood.\textsuperscript{12,14,15} Several theories include delivery by comets or asteroids, the reaction of hydrogen from nebular gas with iron oxides in the Earth’s mantle, and the adsorption of gaseous water onto the accretion disk.\textsuperscript{12,14,15} An accretion disk describes a collection of interstellar grains that aggregate during planet formation,\textsuperscript{12,15,49} and at present, this theory seems the most plausible.\textsuperscript{15} If water adsorbed to these grains, then water could have been delivered to form Earth’s oceans during this process.\textsuperscript{12,15} Moreover, olivine minerals,\textsuperscript{15} such as forsterite,\textsuperscript{12,13} are the most common astronomical minerals, and thus, adsorption of water to forsterite surfaces, such as the (100) and (010) cleavage planes modeled here, has astrobiological implications.

Regardless of which theory for the delivery of water to Earth is true, the adsorption of water onto olivine phases continues to be important throughout Earth’s history. Earth is believed to have formed from water-containing materials,\textsuperscript{16} but debates exist regarding how much water was contained in the mantle throughout history\textsuperscript{18,20} and at present.\textsuperscript{16-20} The amount of water present in the mantle contributes to the existence of plate tectonics on Earth\textsuperscript{17-19} and to the global water cycle,\textsuperscript{18} such as the volumes of Earth’s oceans.\textsuperscript{50} The mineralogic profile of Earth’s mantle is dominated by olivine-rich minerals,\textsuperscript{17,50} and forsterite is the Mg-rich end-member of the olivine series. Therefore, the adsorption of water to forsterite surfaces as well as the functional groups present at the aqueous-mineral interface are important processes to model for understanding the contribution of water to olivine minerals in Earth’s mantle.

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In addition to the presence of forsterite in accretion disks and in Earth’s mantle, this mineral has also been found on the surface of Mars. The perpetuation of this mineral phase throughout the planet’s surface shows that water has not been present for billions of years or that a limited amount once existed because its dissolution rate is relatively rapid with respect to more silica-rich phases.\textsuperscript{21-23} However, there is evidence for water on Mars in the forms of morphological alteration to minerals\textsuperscript{23} and secondary mineral phases that likely resulted from acidic weathering of forsterite on Mars.\textsuperscript{21,23} These investigations into forsterite weathering provide insight into the aqueous alteration history\textsuperscript{22,23} as well as the impact it has on the chemistry of Martian soils.\textsuperscript{21} Forsterite dissolution on Mars is thought have to occurred under a low water/rock ratio,\textsuperscript{21} and in this work, the investigation of the aqueous-water interface for the forsterite (100) surface solvated by 12 H\textsubscript{2}O molecules models such a system.

### 4.4.2 Future Directions

The main focus of the future directions of this work is to simulate the aqueous-mineral interface for the forsterite (100) and (010) cleavage planes. The energy-minimized structure that resulted from the dissociative adsorption of H\textsubscript{2}O molecules onto the forsterite (100) and (010) surfaces will be used as the starting surface. Bulk water will be modeled by adding 1 g/cm\textsuperscript{3} H\textsubscript{2}O molecules to the periodic cell, and the number of H\textsubscript{2}O molecules needed will be determined via the following equations (4 and 5):
\[
\frac{\text{molecules}}{A^3} = \frac{1 \text{g}}{\text{cm}^3} \times \frac{1 \text{cm}^3}{(10^8 \text{ Å})^3} \times \frac{1 \text{mol}}{18.02 \text{g}} \times 6.02 \times 10^{23} \text{molecules per mol}
\]

\[
\frac{\text{molecules in cell}}{A^3} = \frac{\text{molecules}}{A^3} \times (\text{cell}_x \times \text{cell}_y \times \text{cell}_z) A^3
\]

where \text{cell}_x, \text{cell}_y, and \text{cell}_z correspond to the parameters of the vacuum space over the forsterite (100) and (010) slabs. Sixteen \text{H}_2\text{O} molecules are needed for the forsterite (100) cell and 21 \text{H}_2\text{O} molecules for the (010) cell. Inclusion of \text{H}_2\text{O} molecules in the simulation cell according to Equations 4-4–5 models solvation of the surface at the aqueous-mineral interface. The \text{H}_2\text{O} molecules will be energy-minimized separately in a cell whose dimensions were the same as the vacuum portion of the forsterite (100) and (010) simulation cells. DFT-MD simulations using these energy-minimized structures will be run for a 5 ps equilibration period with a timestep of 0.5 fs. Results for structural energy and dynamical analysis will then be collected over an additional 5 ps with a 0.5 fs timestep.

After 10 ps of simulation, the samples will be characterized according to how many of each functional are present on the surface. In addition, the average positions of \text{O} and \text{H} atoms in the simulation cells for the last 5 ps will be analyzed to determine how these atoms are moving during the timescale simulated. The goal of this analysis will be to understand where these atoms are located to provide insight into spectroscopic analysis used to characterize these systems.

Some initial simulations have been performed for the forsterite (100) and (010) surfaces at the aqueous-mineral interface. Twenty-one \text{H}_2\text{O} molecules were used to solvate the forsterite (100) surface while 34 \text{H}_2\text{O} molecules were used to solvate the forsterite (010) surface. As with the monolayer surfaces, a distribution of functional
groups is present on the forsterite (100) and (010) surfaces at the aqueous-mineral interface. In addition, H\(^+\) transfers occur between surface groups and between surface groups and solution, such that the distribution of functional groups present on the surface evolves with time. When H\(^+\) ions adsorb to the surface, the primary sites are Si–O\(^-\), Mg–O\(_{br}\), or Mg–OH, and in the case of Mg–O\(_{br}\) adsorption, this then results in a new reaction site on the surface. H\(_2\)O transfers from solution to surface sites also occur, and these are located at Mg sites on the surface, which has been observed in previous calculations.\(^{38}\)

On the surface of each of these samples at the aqueous-mineral interface, at least one O\(^-\) group is present. One would anticipate that this excess of electron density would be unstable on the surface and readily become protonated. Although H\(^+\) ions are transferred to such sites throughout the simulations, O\(^-\) groups remain. This observation is in accord with previous DFT calculations where a Si–O\(^-\) group was stabilized by solvation by three H\(_2\)O molecules,\(^{52}\) and therefore, such a site can be expected to perpetuate at the aqueous-mineral interface for forsterite.

For these two samples as well as those modeling 1 g/cm\(^3\), an analysis scheme will be designed to capture the events within the simulations that best capture the essential science of the aqueous-mineral interface for forsterite (100) and (010) surfaces. The number of H\(^+\) and H\(_2\)O transfers can be calculated for each solvated forsterite (100) surface and the solvated forsterite (010) surface. The goal is to identify at which functional groups H\(^+\) transfers occur and thus to show which are the more reactive sites on each surface. In addition, H\(^+\) and H\(_2\)O transfers between surface groups and solution also occurred, and these transfers will be quantified and tracked with time. Further, such analysis will provide insight into whether the forsterite (100) or the (010) cleavage plane
is more reactive at the aqueous-mineral interface, particularly since the coverage of these surfaces by H\textsubscript{2}O molecules is very similar among the systems studied.

In addition to quantification of H\textsuperscript{+} and H\textsubscript{2}O transfers, the H-bond lengths can be calculated according to the analysis scheme of Kumar, et al.\textsuperscript{32} The H-bond interactions were shown to be the strongest between Ti–OH sites and H\textsubscript{2}O molecules in a monolayer on the rutile surface, whereas Ti\textsubscript{2}O sites interacted the weakest with H\textsubscript{2}O molecules adsorbed to the rutile surface. Further, the bond lengths on the surface of goethite have been shown to lengthen as additional H\textsubscript{2}O molecules solvate the surface.\textsuperscript{33} A similar approach will be applied here to calculate the bond distances for sites on the forsterite (100) and (010) surfaces.
4.5 References

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Chapter 5

Conclusions and Future Directions

5.1 Summary of Research

The overarching goal of the work presented in this dissertation was to use existing computational tools to model geochemical systems across spatial and temporal scales. Density functional theory (DFT) calculations were employed to model the release of divalent metal ions from individual surface sites on ortho- and framework-silicate mineral surfaces. DFT molecular dynamics (DFT-MD) simulations were used to model the forsterite mineral surface along two different cleavage planes, and these simulations allowed for the description of multiple surface sites simultaneously. Hence, the use of DFT-MD simulations enabled investigation of a larger system size as well as a longer timescale, as bulk physicochemical processes were modeled.

In Chapter 2, Mg\(^{2+}\) release from the forsterite mineral surface from sites of various protonation states was modeled. A reaction mechanism was described for hydrolysis of each type of site, and the barrier heights were also calculated. Ab-initio results were coupled with experimental data, which incorporated the concentration of sites in each protonation state. The mechanisms showed that hydrolysis occurs through two steps, namely the breaking of the Mg–O bond and the incorporation of a H\(_2\)O molecule from the second hydration sphere into the first hydration sphere. In both the gas-phase and aqueous-phase, the protonated sites have the lowest barrier height,
followed by the deprotonated and neutral sites, respectively. In addition, calculations employing aqueous-phase models must be optimized because single-point aqueous-phase calculations yielded unreliable results, and this is particularly important when modeling geochemical processes that occur in aqueous media.

The rate constants for the release of Mg\(^{2+}\) from Mg–O–Si sites were calculated using the barrier heights from the DFT calculations. The rate constants and thus Mg\(^{2+}\) release rates were higher than those measured experimentally,\(^1\)–\(^7\) which was, in part, a result of higher \(E_{app}\) values measured experimentally for the dissolution of forsterite.\(^8\),\(^9\) This shows that the rate-limiting step is not the release of Mg\(^{2+}\) as a result of breaking the last Mg–O bond to the forsterite surface. At the same time, the DFT results corroborate previous experimental and DFT findings. \(^1\)H–\(^{29}\)Si cross-polarization magic angle spinning (CPMAS) NMR experiments showed that a Mg-depleted, Si-rich layer develops on the forsterite surface during dissolution in acidic conditions.\(^10\) When the DFT results from this work are compared with barrier heights for the hydrolysis of Si–O–Si sites on quartz,\(^11\)–\(^14\) Mg\(^{2+}\) ions would be expected to leach first from forsterite during dissolution.

The reaction coordinates were chosen from two possibilities. Initially, the Mg–OH\(_2\) distance seemed reasonable, but decreasing the distance between the H\(_2\)O molecule in the second hydration sphere and the Mg\(^{2+}\) ion did not lead to the breaking of the Mg–O bond connecting the ion with the surface. Instead, such a reaction coordinate led to the exchange of H\(_2\)O molecules around the Mg\(^{2+}\) ion, while the Mg–O bond remained unbroken. Thus, Mg–O bonds on the forsterite surface break as a result of the lengthening of this bond and not from H\(_2\)O exchange around the Mg\(^{2+}\) ion.
In Chapter 3, the link between H₂O exchange reactions and M–O bond break was investigated to determine which of these processes led to release of metal ions to solution. The H₂O exchange reaction around the hexaaqua Mg²⁺ ion was modeled in the gas-phase and aqueous-phase, using both the IEFPCM and super-molecule approximations. The barrier heights using the IEFPCM and super-molecule approaches were comparable to experimental values within computational uncertainty. The high barrier height for this reaction in the gas-phase is likely a result of a high energy TS characterized by distorted pentagonal bipyramidal geometry. In the IEFPCM and super-molecule calculations, however, the TSs had square-pyramidal geometry, indicative of six electron pairs around the Mg²⁺ ion. The different TS geometries for these two phases result in a barrier height for the gas-phase that was ~30–40 kJ/mol higher than the aqueous-phase models. The aqueous-phase model calculations show that this approach is appropriate for these systems.

DFT calculations were used to probe the link between H₂O exchange reactions for hydrated metal ions and dissolution of end-member silicate minerals containing those ions. Protonated Ni–O–Si, Mg–O–Si, and Ca–O–Si sites were found to react via three separate mechanisms. The breaking of the Ni–O bond occurs as a result of approach of a H₂O molecule from the second hydration sphere. For protonated Mg–O–Si sites, approach of a H₂O molecule instead leads to H₂O exchange around the Mg²⁺ ion. Breaking the Mg–O bond is accomplished by lengthening of the bond, in contrast to protonated Ni–O–Si sites. Ca–O–Si sites react via closely coupled reactions for H₂O exchange and hydrolysis. Instead of a traditional H₂O exchange reaction, the second hydration sphere H₂O molecule was absorbed by the Ca²⁺ ion to form a hepta-
coordinated Ca-silicate cluster. Similar to protonated Mg–O–Si sites, the breaking of the Ca–O bond occurred via lengthening this bond, but the precise reaction mechanism was difficult to elucidate.

The barrier heights of these reactions were used to calculate the rate constants for these reactions via the classical TST approximation, and in accord with experimental trends, the rate constants increase in the order Ni$^{2+} < $ Mg$^{2+} < $ Ca$^{2+}$. What is more, rate constants for hydrolysis and H$_2$O exchange of these M–O–Si clusters differ from rate constants for H$_2$O exchange around these ions in solution, and therefore, the Si-containing group as opposed to a sixth H$_2$O molecule contributes to the kinetics of these metal ions. The rate constants for H$_2$O exchange and hydrolysis around protonated Ni–O–Si and Mg–O–Si sites are lower than those for Ni(H$_2$O)$_6^{2+}$ and Mg(H$_2$O)$_6^{2+}$. For protonated Ca–O–Si sites, however, the rate constant for Ca(H$_2$O)$_6^{2+}$ is lower than those for the formation of the hepta-coordinated complex and hydrolysis.

As with the calculations in Chapter 2, there were two possibilities for the reaction coordinate. Initially, the distance between the metal ion and the H$_2$O molecule in the second hydration sphere was used. If a H$_2$O exchange reaction occurred and not hydrolysis, then the M–O bond was lengthened until it broke. A second PES scan was performed to model the absorption of the second hydration sphere H$_2$O molecule into the first. In this way, the correlation between H$_2$O exchange reaction rates and hydrolysis rates for protonated sites on end-member silicates was modeled.

In Chapter 4, DFT-MD calculations via the VASP were used to simulate the adsorption of a monolayer of H$_2$O molecules to the forsterite (100) and (010) cleavage planes. Both the associative and dissociative adsorption processes were modeled for each
cleavage plane, and the energy-minimized surfaces were characterized by a distribution of functional groups that included O\textsubscript{br}, O\textsuperscript{-}, OH, and H\textsubscript{2}O sites regardless of whether the starting surface was covered with dissociated or intact H\textsubscript{2}O molecules. For both the (100) and (010) surfaces, dissociative adsorption is predicted to be a lower-energy process.

In addition to investigating the surface structure for the adsorption of H\textsubscript{2}O molecules, a comparison was made regarding the stability of the (100) and (010) cleavage planes for forsterite, and the (100) plane was determined to be lower in energy. This is in accord with a previous study\textsuperscript{19} but in contrast to another.\textsuperscript{20} However, care was taken here to cut the slabs such that dangling bonds were minimized, and thus the formation of defects on the surface was avoided. The presence of both dangling bonds and surface defects contributed to the instability of forsterite (100) in a previous study, which is likely why it was found to be higher in energy than the (010) cleavage plane.\textsuperscript{20}

5.2 Expected Implications

There were several goals to the work described in this dissertation. The first was to use existing computational tools to model geochemical systems across spatial and temporal scales. The second was to provide molecular scale insight into experimental observations. The third goal of this work was to combine experimental data with calculated results. The fourth goal was to initiate a study of the forsterite surface at the aqueous-mineral interface so that larger-scale investigations of the forsterite surface at
various stages during dissolution could be modeled, and these future calculations are described in Section 5.3.

These goals were each successfully attained. Two different timescales and spatial scales were modeled through metal release from single surface sites on silicate surfaces as well as by simulating the aqueous-mineral interface on a unit cell sized scale for forsterite. Molecular scale insight was provided by determining how the final M–O bond connecting the metal surface breaks as a result of H₂O approach for protonated Ni–O–Si sites and as a result of M–O bond lengthening for protonated Ca–O–Si sites and for Mg–O–Si sites in various protonation states. In addition, differences in calculated Mg²⁺ release rates from forsterite as well as surface characterization studies for unit cell sized forsterite systems show that rate-controlling dissolution reactions likely occur at specific reactive sites on the surface versus all sites present. Experimental data were combined with calculated results in Chapter 2, where the coverage of the forsterite surface by H⁺ or OH⁻ ions measured experimentally versus pH was combined with ab-initio results to calculate a rate for Mg²⁺ release. Also in Chapter 3, the rate constants calculated for H₂O exchange and hydrolysis for protonated M–O–Si (M = Ni²⁺, Mg²⁺, and Ca²⁺) clusters are compared with those measure for H₂O exchange reactions around these ions in solution. The initial work for simulating the aqueous-mineral interface for two forsterite cleavage planes was begun. Initial findings showed that the (100) cleavage plane is lower in energy than the (010) plane and that dissociative adsorption of H₂O molecules is likely the lower energy process. However, for both associative and dissociative adsorption, the energy-minimized surfaces are characterized by sites with a distribution of functional groups.
The implications of this work pertain to integrating computational tools into geochemical kinetics investigations. Existing computational technology allows for detailed descriptions of the mechanisms throughout which geochemical species react. Further, the employment of multi-scale tools allows for insight into which processes are rate-controlling and if any act concertedly. Examples include the coupled H$_2$O exchange and hydrolysis reaction for protonated Ca–O–Si and the presence of a distribution of functional groups on the forsterite surface. Lastly, such a multi-faceted approach begs to question whether these approaches can be similarly adapted for other minerals. In fact, studies for quartz, aluminosilicates, and rutile have already been performed, and possibilities for future study include mixed-cation silicate minerals.

In addition, the description of molecular scale models provides insight into experimental observations of geochemical systems on much larger temporal and spatial scales. For example, current experimental studies use laboratory-scale models to replicate field conditions and isolate which are the controlling factors to dissolution rates measured in the field, and reasons for discrepancies between the dissolution rates for minerals at the different time and spatial scales are suggested. White and Brantley showed that the surface roughness in part contributed to differences between lab and field dissolution rates, while Navarre-Sitchler and Brantley showed that a number of contributing factors including reaction mechanism account for the discrepancies in dissolution rates between scales. In this way, molecular scale understanding of their geochemical systems would further contribute to their conclusions. For example, molecular scale models enable the amount of energy to be calculated for simplified systems where the most fundamental phenomena can be simulated. Therefore, the
aspects of the system which are the energetically controlling the observed process can be identified, and in these experimental examples,\textsuperscript{31,32} such molecular scale insight would explain why two different minerals within the same sample weather at different rates.

5.3 Future Directions

Studies of the aqueous-mineral interfaces for the forsterite (100) and (010) surfaces are currently in progress. The number of H\textsubscript{2}O molecules required to simulate the presence of bulk water over the forsterite (100) and (010) slabs is being energy-minimized in the vacuum space above the slab, and once finished, these molecules will be added to the simulation cell. DFT-MD simulations will be performed for 10 ps, where a 5 ps equilibrium period will be followed by 5 ps of analysis time. The location of O and H atoms within the simulation cell will be analyzed with time, and the number of H\textsuperscript{+} and H\textsubscript{2}O transfers over time will be calculated. Both H\textsuperscript{+} and H\textsubscript{2}O transfers are anticipated to occur between groups on the surface and between surface groups and species in solution. This would lead to a change in the number of functional groups on the surface throughout the surface, but a distribution of functional groups is expected to remain throughout these simulations. In addition, the O–H bond lengths and H-bond interaction distances will be calculated so that these quantities can be compared to experimental measurements. These distances were found to change from equilibrium at the aqueous-mineral interface for rutile,\textsuperscript{29} and thus a similar analysis will be conducted here.
Moreover, the studies described in Chapters 2–4 focused on pristine mineral surfaces. During dissolution, however, the mineral surface is physically as well as chemically altered, and for forsterite, a Mg-depleted layer develops. At this point, no molecular scale description of such a layer has been delineated, but this layer is known to consist of Si atoms with three Si–O–Si bonds. Thus one possible future direction of this work is to employ DFT-MD simulations in order to investigate the stability of a forsterite surface where Mg$^{2+}$ ions have been removed and isolated SiO$_4^{4-}$ tetrahedra remain.

Similarly, the presence of re-polymerized SiO$_4^{4-}$ tetrahedra has been characterized via $^1$H-$^{29}$Si cross-polarization magic angle spinning (CPMAS) NMR, but the mechanism through which the surface reconnects these isolated tetrahedra is not yet known. This re-polymerization of silica species on the forsterite surface could be investigated with a multi-scale approach similar to the one adopted in this dissertation. For example, molecular clusters comprised of several Mg–O–Si linkages could be designed and then Mg$^{2+}$ ions removed, leaving behind SiO$_4^{4-}$ tetrahedra. The energy for this process could be calculated. The ensuing Mg-depleted clusters could be reacted with H$_2$O molecules to show how these tetrahedra connect to form the silica-rich layer observed experimentally. In addition, a forsterite sample comprised of several unit cells could be modeled, and again Mg$^{2+}$ ions removed. Here, DFT-MD simulations modeling this Mg-depleted forsterite surface at the aqueous-interface could be performed such that H$^+$ and H$_2$O from solution, for example, would be available to react with the surface, and surface reconstruction to yield a silica-rich layer could be modeled over time.
Once these Mg-depleted and re-polymerized samples have been sufficiently energy-minimized, they would be used to calculated NMR chemical shifts. Davis et al.\textsuperscript{10} observed chemical shifts at –62 ppm for the pristine forsterite surface, corresponding to $Q^0$ Si atoms – that is, those Si atoms without any Si–O–Si bonds. At 190 h of dissolution in acidic media, a peak at –93 ppm, corresponding to $Q^3$ Si atoms – those Si atoms with three Si–O–Si bonds – began to develop. The molecular clusters representing intact forsterite and Mg-depleted forsterite surfaces could be used in an attempt to replicate the experiments of Davis \textit{et al.}

The expected implications of these future directions are to continue to develop a molecular scale picture of forsterite dissolution and to discern an appropriate method for calculating NMR shifts of minerals. The mechanisms for Mg$^{2+}$ release from sites of various protonation states and for H$_2$O exchange and hydrolysis of protonated M–O–Si (M = Ni$^{2+}$, Mg$^{2+}$, and Ca$^{2+}$) sites were described in Chapters 2 and 3, respectively. However, questions remain as to the stability of the surface after Mg$^{2+}$ release and the process through which SiO$_4^{4–}$ tetrahedra re-polymerize. Previous work where NMR chemical shifts were calculated showed that these results are heavily dependent upon which functional and basis set were chosen.\textsuperscript{37-39} Therefore, determining the NMR chemical shifts for these geochemical systems will also allow for a more systematic approach of calculating NMR chemical shifts with ab-initio methods.
5.4 References


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