ELASTIC PROPERTIES OF CEMENT PHASES USING
MOLECULAR DYNAMIC SIMULATION

by

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AUTHOR’S DECLARATION

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ELASTIC PROPERTIES OF CEMENT PHASES USING
MOLECULAR DYNAMIC SIMULATION

Mohamed Arar, Master of Applied Science, Civil Engineering, Ryerson University, 2016

Abstract

The goal of this thesis is to shed light on the elastic properties, especially the Young modulus, of clinker phases and hydration products of cement paste through molecular dynamic simulation by COMPASS force field. The parameters that can affect the elastic properties of cement phases were also targeted, with special attention paid to analog C-S-H minerals, in which the Tobermorite family and Jennite were simulated to render their structures representative of C-S-H structure.

The molecular dynamic simulations of this study show that CO force field can be applicable for most clinker phases and hydration products. Jennite, with its porosity and finite silicate chain accounted for, can be considered the closer and representative structure of C-S-H. In addition, this study confirms the important effect of C/S ratio, packing factor and chain length on elastic properties of C-S-H.
Acknowledgements

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Dedication

This thesis is dedicated to the memory of my father. It is also dedicated to my mother, my wife, my wonderful son and two daughters, and to all my family who supported and helped me accomplish my goal in attaining higher education.
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Chapter 1  Introduction

Over the past few years, there has been a growing interest and focus on sustainable buildings. Materials are now evaluated for environmental impact associated with their manufacturing and transportation. Concrete is the most commonly used construction material, with a 4% annual increase in cement production due to demand (Kusuma, Budidarmawan, & Susilowati, 2015). However, the construction of city infrastructure, which is the base for economic growth and job creation, results in a release of significant amounts of CO₂ and greenhouse gas emissions. One way to decrease environmental impact and contribute to sustainable development is to improve concrete quality, in which cement is the primary ingredient. Improved concrete quality increases the lifespan of structures, reducing environmental impact and project cost (Kusuma et al., 2015). For instance, improved concrete quality with low standard deviation (S=2.4MPa) has a 41% smaller carbon footprint than low quality and high standard deviation (S=8.6MPa) concrete (Kusuma, et al., 2015).

Nanotechnology can be used to develop a new material with higher performance and sustainability. Nanotechnology was introduced by physicist Richard Feynman in his 1959 lecture, (Sanchez & Sobolev, 2010) in which he suggested the possibility of manipulating atoms and molecules to create new materials with improved properties. Although nanotechnology has different meanings from country to country, all definitions agree that it deals with elements at a very small scale, generally less than 100 nm (Sanchez & Sobolev, 2010). The concept of nanotechnology was well developed in the 1990s in chemistry, biology, physics and other fields. It has two approaches: top down and bottom up. In the top down approach, bulk material is reduced to nanoscale level without changing the properties of the material. The purpose of this approach is to create small devices and materials at
nanoscale level such as computer chips, nano silica, carbon nanotubes, nano wire and nano powder. In the bottom up approach – also called “molecular nanotechnology” (Sanchez & Sobolev, 2010) – the material is made by assembling atoms and molecules. Although molecular nanotechnology has made advances in fields such as medicine, energy, biotechnology and information technology, it is still not widely used in cementitious material. The major progress in its use in cement paste has been to understand the structure and mechanical properties of key cement paste components, as well as the source of forces between particles.

The binding element in concrete is the Calcium Silicate Hydrate (C-S-H). When seen from the bottom up approach, it is composed of atoms that bind together through different types of interaction such as chemical bond, Coulombic, van der Waal interactions and others. The purpose of the bottom up approach, which deals with manipulation of the structure at nano scale, is to develop a new generation of materials with high mechanical properties, durability, self-sensing, self-healing and self-cleaning properties.

The purpose of computation is to predict the behavior and the properties of the material in different conditions such as high temperature (melting), high pressure and others, with the use of physics and chemistry theories and principles.

Material computation is also helpful when the experimental and computational results support each other. Results of computational methods can help predict the behavior and engineering properties of cement paste and aid in the development of new materials. It is also beneficial because it reduces the number of experiments needed when developing a new material, and its results are used for comparison with experimental results.

There are two main approaches in material computation to model interaction between atoms: the empirical approach and quantum mechanics (ab initio). In the empirical approach, a set of simple equations (force field) describes the interactions between atoms. The quantum mechanics approach is
based on the solution of the Schrödinger wave equations of all electrons in the system. Four methods are derived from these two approaches: lattice energy modeling, calculation of lattice dynamics, Monte Carlo simulation and molecular dynamic simulation (MD).

Lattice energy modeling, calculation of lattice dynamics and Monte Carlo simulation all require time-consuming, expensive and rigorous computation with very powerful computers. Therefore, for this thesis, an empirical approach using MD simulation was chosen to speed simulation time, better understand the structure of materials at nanoscale level, predict and improve its properties at micro/macro scale level. Many efforts have been made to determine the structure of calcium silicate hydrate. However, because C-S-H has an amorphous structure that changes over time, its structure is not yet defined, which is the major challenge in cement paste modeling. Experimental methods using NMR, XANS, nano indentation and other tools have provided some information about the structure of C-S-H; it is a layered structure composed of short-range crystals. However, some parameters can’t be determined experimentally. The quantum mechanics computational method (especially the ab-initio method) offered additional information, including the cohesion forces between C-S-H and the relation between the C-S-H structure and Tobermorite family and Jennite, which cannot be determined by experimental tools.

In addition to the similarity with Tobermorite and Jennite, Pellenq et al. (2009) proposed a C-S-H model that can bridge the gap between theoretical and experimental results. However, the proposed structure is not well developed and has limitations. Shahsavari et al. (2009) carried out a dynamic simulation using Density Function Theory (DFT) method using Tobermorite 9A, 11A and 14A. Those authors calculated a Young modulus of 95.06 GPa, 82.82GPa and 51.9 GPa for Tobermorite 9A, Tobermorite 11A and Tobermorite 14A, respectively. Dolado et al. (2007) studied the polymerization of the silicate \((\text{SiOH})_4\) in the presence of ions. They concluded that increased calcium content (Ca/Si) slows the polymerization of the silicate chain. Quantum mechanics methods such as ab initio and DFT are complex, expensive and
time-consuming methods, and require robust computers, which make these methods difficult and reveal the need for further comprehensive and simplified studies on C-S-H simulation.

Using less complicated methods with regular computers and reduced simulation time to achieve reliable results is the principal goal of this study. This can be achieved by using molecular dynamic simulation based on Newton law. The force field used in the molecular dynamic simulation can be considered as the principal parameter that affect the results of the simulation. Among the force fields incorporated in the Materials Studio software (Materials Studio v7), COMPASS force field is an option for reliable results. COMPASS force field has been broadly used for different ranges of molecules and polymers. However, for cement paste components, and especially C-S-H, it has only been used in limited studies. In this study, COMPASS force field is used to determine the elastic properties of clinker phases and cement hydration products such as Portlandite, Ettringite and C-S-H. Special attention was given to C-S-H and its analog minerals including the Tobermorite family and Jennite.

Elastic properties are the main parameters that control the short-term deformation of any structure. For this reason, this research targeted elastic properties such as bulk modulus, shear modulus, Poisson ratio and Young modulus in the dynamic simulation. However, because of the direct relationship between the Young modulus and other elastic properties, most discussions of simulation results of analog C-S-H minerals are based on Young modulus values, as are discussions of the influence of different parameters (density, packing factor, W/Ca, W/Si, porosity) on the elastic properties of analog C-S-H minerals, and the comparison between the elastic properties in this study and those mentioned in previous literature.
**Thesis outline:**

**Chapter 2** covers the basic chemistry of cement and main clinker phases including C₃S, C₂S, C₃A and C₄AF. Hydration process and mechanism, the multiscale level of cement paste and the proposed models for C-S-H gel at its macro-, micro-, nano-scale and atomic-scale levels are also presented.

**Chapter 3** is devoted to the computational method used in this thesis to predict the mechanical properties of the different phases of cement by using molecular mechanics to calculate the energy potential in the system. The chapter also covers the steps required for molecular modelling, including energy minimization, dynamic simulation and production run.

**Chapter 4** deals with the results of the molecular dynamic simulation by Compass force fields of C₃S, C₂S, C₃A and C₄AF. The elastic properties of the clinker phases resulting from this study are also compared to the experimental results, and the effect of packing density on elastic modulus is discussed.

**Chapter 5** covers the elastic properties of Portlandite and Ettringite, the major hydration products of cement reaction. It also includes a comparison between the results of this study and those in literature using other experimental and numerical methods.

**Chapter 6** is devoted to the calculation of the elastic properties of analog C-S-H minerals such as the Tobermorite family and Jennite. The effects of C/S ratio, density and water on elastic modulus of analog C-S-H minerals are also discussed.

**Chapter 7** is focused on the results of two tested methods to reduce the Young modulus of Tobermorite 14A and Jennite to be closer to that of C-S-H. The effects of C/S ratio, silicate chain and porosity on Young modulus results are also discussed.

**Chapter 8** summarizes the general conclusions of this study and offers recommendations for future research.
Chapter 2  Chemistry of Portland cement

2.1 Introduction

Cement is a substance used to bind sand and aggregates. The most commonly used variety is Portland cement; when mixed with water, chemical reactions occur between the components of cement and water to produce a hardened paste. The reaction that causes the setting and hardening of the paste is called a hydration reaction. The Egyptians were the first to use gypsum mortar – made from the calcination of impure gypsum at 130°C (Mindess and Young, 1981) – as binding material. When gypsum is mixed with a small quantity of water, it sets and hardens. However, it is a soluble mortar and cannot harden under water. Lime mortar was also used by the Egyptians; it was prepared by calcining limestone at 1000°C (Mindess et al., 2003). Greeks and Romans also used volcanic ash (Pozzolan). When ground and mixed with lime, sand and water, it creates a mortar that is resistant to water when set and hardened. However, most Romans knowledge – such as burning lime and adding Pozzolan – was lost during the Middle Ages. It was not until 1756, when John Smeaton conducted experiments with lime and pozzolan, he found that the best limestone is one containing a high proportion of clay mineral. Another remarkable discovery was made by Joseph Aspdin, who patented Portland cement in 1824; it is prepared by calcining and mixing ground limestone with fine clay in a kiln (Mindess et al., 2003) then finely grinding it to be used as cement.
2.2 Clinker Phases

2.2.1 Alite, C₃S

Alite (Ca₃SiO₅) is the impure form of C₃S. It is a highly reactive compound because of its high calcium content, and a major constituent of Portland cement clinker, with a range of 50 to 90% of clinker phase (Barnes and Bensted, 1983). Tricalcium silicate is a stable polymorph at temperatures between 1250° - 2150°C. It becomes unstable at temperatures lower than 1250°C but preserves its stability state with rapid cooling. Slow cooling can change Alite structure to Belite or lime (CaO) (Barnes and Bensted, 1983). As the temperature in the clinker changes, Alite undergoes different states as shown:

\[
R_{1060^\circ C} \leftrightarrow M_{1050^\circ C} \leftrightarrow M_{900^\circ C} \leftrightarrow M_{980^\circ C} \leftrightarrow T_{920^\circ C} \leftrightarrow T_{600^\circ C} \leftrightarrow T_1
\]

(T: Triclinic, M: Monoclinic and R: rhombohedral)

The crystal structures of these phases are similar with small distortion in SiO₄ tetrahedral. C₃S has a crystalline structure composed of Ca²⁺, SiO₄²⁻, O²⁻ ions with a C/S ratio of 3:1. Figure 1 shows the unit cell of triclinic C₃S.

![Unit cell of C₃S](image)

Figure 1: Unit cell of C₃S where Ca is presented in green, silica in orange, oxygen in red, and the white line represents the unit cell.

In industrial clinker, C₃S is mixed with foreign ions such as Mg²⁺, Al³⁺ and Fe³⁺, which are present in the raw mix. These impurities prevent the conversion of C₃S to T₁, which is an unstable phase, and
stabilize at high temperature such as M₁ or M₃. These impurities also prevent Alite from decomposing into dicalcium silicate C₂S or lime (CaO) (Barnes and Bensted, 1983). The X-ray diffraction of triclinic C₃S is shown in Figure 2.

![X-ray powder data (CuKα) diagrams for triclinic of C₃S](image)

**Figure 2: X-ray powder data (CuKα) diagrams for triclinic of C₃S**

### 2.2.2 Belite, C₂S

Belite (Ca₂SiO₄), the impure form of C₂S, is the second important clinker phase. It is also the largest, constituting 10 to 40% of the total clinker phase (Barnes and Bensted, 1983). Its hydration provides a similar strength as Alite, with a long period of hydration. It can be found in five types of polymorphs (α, α’, α’, H, α’, L, β and γ), which differ in terms of structure and hydraulic properties. During cooling, α-C₂S goes through these different polymorphs before transforming to β-C₂S at 630°C. The α’₁_H polymorph is the more hydraulic form of Belite; in contrast, the γ polymorph is not reactive which can cause the disintegration (dusting) and cracking of the clinker (Barnes and Bensted, 1983). If the β-C₂S is not stabilized during cooling by the inclusion of impurities, γ –C₂S is less dense and can cause cracking. The β-C₂S can be stabilized at room temperature by doping impurities such as boric oxide, phosphorus pentoxide and magnesia. Belite can be formed in a number of ways, but is primarily formed with the reaction of lime and silicon. However, it can also be formed from Alite (Barnes and Bensted, 1983).
Chemistry of Portland cement

\[ C_3S \leftrightarrow C_2S + C \] (Barnes & Bensted, 1983)

H: high temperature; L: low temperature

The crystal structure of all types of Belite is composed of SiO$_4^{4-}$ tetrahedrals and Ca$^{2+}$. The arrangements of these ions are similar, with some differences in the $\gamma$-C$_2S$ (Taylor, 1997). The C$_2S$ structure differs from Alite by the absence of O$^{2-}$. Figures 3 and 4 show the unit cell of monoclinic $\beta$-C$_2S$ and the X-ray diffraction of $\beta$-C$_2S$.

![Figure 3: Unit cell of C$_2S$ where Ca is presented in green, silica in orange, oxygen in red, and the white line represents the unit cell](image)

![Figure 4: X-ray powder data (CuKα) diagrams for monoclinic of $\beta$-C$_2S$](image)

2.2.3 Tricalcium aluminate, C$_3$A

The third important component in ordinary Portland cement is tricalcium aluminate (Ca$_3$Al$_2$O$_3$ or C$_3$A), which represents 10% of the clinker phase. Tricalcium aluminate does not appear in pure form, but rather, it is modified by the substitution of Al and Ca with Na$^+$ and Si$^{4+}$, Fe$^{3+}$ and Mg$^{2+}$ or K$^+$. The
structure of tricalcium aluminate has a cubic form. However, due to the high quantity of alkalis in raw clinker, the C₃A can have an orthorhombic or monoclinic structure. The structure of C₃A is composed of rings of six AlO₄ tetrahedrals and Ca²⁺ ions. The reactivity of these polymorphs depends on the quality of foreign ions existing in the lattice. Pure C₃A contains 62.3% CaO and 37.7% Al₂O₃. The total replacement of foreign ions is around 13% for the cubic and 20% for the orthorhombic (Taylor, 1997). The unit cell of C₃A and the X-ray diffraction of C₃A are shown in Figures 5 and 6, respectively.

Figure 5: Unit cell of C₃A where Ca is presented in green, aluminum in magenta, oxygen in red, and the white line represents the unit cell

Figure 6: X-ray powder data (CuKα) diagrams for cubic of C₃A Ferrite C₄AF

2.2.4 Ferrite C₄AF

The chemical formula of calcium aluminate ferrite can be written as C₂(A,F). The structure of the ferrite phase is composed of Ca²⁺ ions surrounded by several oxygen ions, with Al³⁺ and Fe³⁺
distributed between octahedral and tetrahedral sites. In the ferrite phase, the $F^{3+}$ ions may be substituted by $Mg^{2+}$, $Si^{4+}$ and $Ti^{4+}$. Figures 7 and 8 show the unit structure on elevation (3D view) and the X-ray diffraction of the ferrite phase ($C_4AF$).

![Figure 7: Unit cell of $C_4AF$ where Ca is presented in green, aluminum in magenta, oxygen in red, iron in violet, and the white line represents the unit cell](image)

![Figure 8: X-ray powder data (CuKα) diagrams for triclinic of $C_4AF$](image)
2.3 Hydration of Portland cement

Hydration is the reaction of non-hydrated cement with water that results in a strong, hard material called paste. It involves a number of reactions which take place simultaneously or one after another. The rate of hydration is affected by a number of parameters such as the composition of the clinker (C₃S, C₂S, C₃A, C₄AF…), presence of foreign ions in the clinker, fineness of the cement particles (surface area), w/c ratio, curing temperature and the presence of chemical admixture and supplementary cementitious material (SCM). Due to the complexity of the hydration reaction in Portland cement, it is preferable to discuss the hydration of the individual clinker phase separately.

2.3.1 Tricalcium silicate (Alite)

The hydration products of C₃S at ambient temperature are an amorphous calcium silicate called C-S-H and calcium hydroxide Ca(OH)₂. The hydration of Alite controls the setting and early strength of Portland cement paste. About 70% of the C₃S reacts in 28 days and 90% will hydrate within one year (Taylor, 1997). The hydration reaction of C₃S can be written as:

\[
2\text{C}_3\text{S} + 6\text{H} \rightarrow \text{C}_3\text{S}_2\text{H}_3 \text{(gel)} + 3 \text{CH}
\]

Tricalcium silicate   Water   C-S-H   Calcium hydroxide

The formula of C₃S₂H₃ is only an approximation because the hydration product varies over a wide range. In addition to the formation of C-S-H, calcium hydroxide is also formed because the ratio of C/S is lower in C-S-H than in the original C₃S, which is 3:1 (Mindess et al., 2003).

2.3.2 Dicalcium silicate (Alite)

The hydration rate of C₂S is lower than C₃S because C₂S ions are denser than those in C₃S, which attracts less water for hydration (Odler, 2000). When C₂S is mixed with water at room temperature, full hydration takes several years. The hydration reaction of C₂S can be written as:

\[
2\text{C}_2\text{S} + 4\text{H} \rightarrow \text{C}_3\text{S}_2\text{H}_3 \text{(gel)} + \text{CH}
\]
The hydration product of C\textsubscript{2}S is almost identical to that of C\textsubscript{3}S, with a smaller amount of calcium hydroxide (CH).

### 2.3.3 Tricalcium aluminate (C\textsubscript{3}A)

When mixed with water, tricalcium aluminate reacts very quickly to form intermediate phases. The hydration products of C\textsubscript{3}A are C\textsubscript{2}AH\textsubscript{8} and C\textsubscript{4}AH\textsubscript{19} by the following reactions:

\[ 2C3A + 27H \rightarrow C2AH8 + C4AH19 \]
\[ C3A + CH + 19H \rightarrow C4AH19 \]

C\textsubscript{2}AH\textsubscript{8} and C\textsubscript{4}AH\textsubscript{19} belong to the group called AFm (alumina, ferric, monosulAFte). Both C\textsubscript{4}AH\textsubscript{19} and C\textsubscript{2}AH\textsubscript{8} are thermodynamically unstable and convert to C\textsubscript{3}AH\textsubscript{6}, which belong to the hydrogarnet group.

\[ C4AH19 + C2AH8 \rightarrow 2C3AH6 + 15H \]

C\textsubscript{4}AH\textsubscript{19} forms in Portland cement only when the calcium sulfate has been consumed. C\textsubscript{3}A reacts with the calcium sulfate to form a tricalcium aluminate trisulfate hydrate called Ettringite. After the consumption of calcium sulfate, the Ettringite reacts with the not-yet-consumed tricalcium aluminate to form tricalcium monosulfate (C\textsubscript{3}A\textsubscript{C}\textsubscript{S}\textsubscript{12}H or C\textsubscript{4}A\textsubscript{S}\textsubscript{H}\textsubscript{12}). The reaction of C\textsubscript{3}A in the absence of gypsum leads to so-called “flash setting,” an undesirable phenomenon that reduces workability and final strength. Gypsum helps regulate C\textsubscript{3}A hydration to avoid flash setting. The presence of C\textsubscript{3}A in cement has a negative effect on concrete durability because of the presence of AFm phases, which, when exposed to sulfate solution, can cause sulfate attack (Mindess et al., 2003). In addition, delayed Ettringite (DEF) occurs when concrete is subject to high temperatures (70-88°C) during the early age of curing (Mindess et al., 2003). High temperature curing destroys the originally formed Ettringite. After cooling, the sulfate is available to form Ettringite again, which can cause expansion in hardened concrete.
2.3.4 Aluminoferrite (C₄AF)

The hydration products of the ferrite phase are similar to those formed during the hydration of C₃A; they contain Fe³⁺ as well as Al³⁺. The hydration reaction is slower than in C₃A, and it decreases as iron content increases. The hydration reaction of C₄AF produces less heat in comparison with C₃A. High C₄AF content in the cement results in better resistance to sulfate attack (Mindess et al., 2003).

2.4 Mechanism of hydration

Different stages can be observed during the hydration of cement. These stages are monitored by heat liberation during hydration:

2.4.1 Pre-induction period (Stage I)

As soon as the cement comes in contact with water, rapid dissolution of ionic species into the liquid phase occurs and the first C-S-H forms. The dissolution of alkalis and sulfate present in the cement occurs very quickly, which leads to K⁺, Na⁺ and SO₄²⁻ ions in the liquid solution. The dissolution of C₃S increases the concentration of Ca²⁺ and OH⁻ in the liquid phase:

\[ \text{C₃S} + \text{water} \rightarrow 3\text{Ca}^{2+} + 4\text{OH}^- + \text{H}_2\text{SO}_4^{2-} \]

C₃A dissolves and reacts with Ca²⁺ and SO₄²⁻ in the liquid phase to form Ettringite. The hydration of C₃A in the pre-induction period varies from 5 to 25% (Lea and Hewlett, 1998). The ferrite phase reacts in a similar manner as C₃A, leading to the formation of the AFt phase. The β-C₂S reacts with water and leads to the formation of C-S-H and the dissolution of Ca²⁺ and OH⁻ in the liquid, which increase the liquid concentration. Early hydration is slowed due to hydration product being deposited onto the cement grain, forming a barrier between the non-hydrated material and the solution. This barrier causes increased ion concentration due to the dissolved elements in the liquid phase.

2.4.2 Induction (dormant) period (Stage II)

The hydration rate in this period slows for few hours, causing the Portland cement concrete to remain plastic during this period. The causes of the induction period have been the subject of many theories, including: 1) the theory that a protective layer is formed around the C₃S grains and the induction
period ends only when this layer is destroyed or becomes more permeable (Jennings, 1986), and 2) that the rate of the reaction in the nucleation period is controlled by the concentration of ions in the solution. When the concentration of Ca\(^{2+}\) and OH\(^{-}\) reaches a critical value, hydration products such CH and C-S-H start to crystallize (nucleation) (Wu and Young, 1984). CH crystals from the solution and C-S-H form on the surface of the grains. The induction period ends when crystal growth begins. Figure 9 shows the different phases of cement hydration: dissolution, nucleation and crystallization of CH and C-S-H.

Figure 9: Top figure shows concentration of Ca\(^{2+}\) in solution with time; bottom figure shows different stages of hydration as a function of time (Barnes, 1983)

2.4.3 Acceleration period (Stage III)

This phase occurs in the first 3-12 hours after mixing. An increase in the rate of hydration occurs in this phase, with accelerated dissolution of calcium silicate and aluminate as a consequence increased liberated heat. The crystal CH (Portlandite) precipitates from the liquid phase with a decreased concentration of Ca\(^{2+}\) and OH\(^{-}\) in the pore solution. The calcium sulfate continues to dissolve in the liquid and SO\(_4^{2-}\) concentration decreases due to the formation of Ettringite, and also due to adsorption of SiO\(_4^{2-}\) for the formation of C-S-H (Lea and Hewlett, 1998).
2.4.4 Post-acceleration stage (Stage IV)

In this phase, the rate of hydration decreases slowly and hydration becomes diffusion-controlled. C-S-H continues to form from C\(_3\)S and C\(_2\)S, with the contribution of C\(_2\)S becoming more pronounced with time. When the calcium sulfate is consumed, SiO\(_4^{2-}\) concentration is reduced. The AFt phase reacts with additional C\(_3\)A and C\(_2\)(A,F) to form monosulfate aluminate. With a high w/c ratio, the hydration process continues until all the anhydrite cement is consumed. Larger cement particles may take longer to hydrate. At low w/c, the hydration reaction may stop, even with the availability of significant amounts of non-reacted cement.

2.5 Multiscale level of cement paste

Predicting properties of cement paste or concrete is difficult due to the composite behavior, heterogeneity of the material and the existence of multi-scale length. An understanding of multi-scale levels of cement paste and the interactions between these scales is fundamental to understanding the properties of concrete. Concrete scale can be divided in three levels: macro, micro and nano, as illustrated in Figure 10.

![Figure 10: Structural hierarchy in concrete](image)

(a) Optical image of concrete showing aggregates in a matrix of hardened cement paste, (b) Back scattered electron image of hardened cement paste showing unhydrated Ca\(_5\)SiO\(_5\) (brightest phase),
Ca(OH)₂ (dark gray), C₃S-H (light gray), and apillary pores (black), (c) Agglomerated particles of C-S-H creating a gel-like structure of high surface area, (d) Layered crystal structure of 1.4-nm tobermorite, a natural mineral whose structure is related to that of C₃S-H at the sub-particle level.

2.5.1 Macro scale

The term macro-scale refers to gross microstructure visible to the human eye (1/5 mm) (Mehta and Monteiro, 2014). At macro-scale level, structures have two phases: a composite of aggregate particles of different sizes and shapes and the binding element composed of hydrated cement paste. Macro pores (entrapped air) can also be seen, having been caused due to a lack of vibration, which has a negative effect on the mechanical properties of the paste.

2.5.2 Micro structure scale

The microstructure is the 10⁶ magnified portion of the macrostructure. At the microscopic level, the two phases are not homogenously distributed and each scale represents a new composite. For example, the hydrated cement appears to be dense in some areas and porous in others. In addition, the hydrated cement paste appears to be different close to the aggregate zone (ITZ: interfacial transition zone). C₃S-H at micro-scale depends on many factors including the nature of the cement, type, admixture, temperature, period of hydration and w/c ratio. Study of the microstructure includes morphology (sizes and shapes), surface area, density, porosity, pore shape and pore distribution (Ramachandran, 1984). Many parameters at the microstructure level are connected to each other; no individual parameter can control the properties of the paste. The morphology of C₃S-H at micro-scale level has been described as foils and flakes, crumpled sheets and rods, and tube for AFt, depending of the degree of hydration (Ramachandran, 1984). At a later age, two types of products in C₃S-H can be distinguished: the “inner product” formed in the boundaries of the original anhydrous grains and the “outer product” filling the space originally filled with water. Diamond (2004) found that the ratio of Ca/Si is higher in the inner products than in the outer ones.
In addition to the solid phase, the hydrated cement paste contains capillary voids, which are the empty spaces that occur when the hydration product takes the space of water and cement. Capillary voids range from 3-5 μm at early ages to 10-50 nm at late ages (Mehta and Monteiro, 2001). As cement hydration occurs, pore volume decreases, and the channels connecting them are broken. Also, the higher the w/c, the higher the pore volume.

At this scale, calcium hydroxide (CH) can be distinguished from C-S-H because it has a different size and shape. The morphology of CH is affected by available growth space, temperature and impurities in the cement (Mehta and Monteiro, 2001).

2.5.3 Nano scale level

At the nano-scale level of less than 10^{-9} m, a number of methods are used to study the structure of C-S-H: nuclear magnetic resonance (NMR), scanning electron microscope (SEM), transmission electron microscopy (TEM), atomic force microscopy (AFM), small-angle scattering (SANS) and small-angle X-ray scattering (SAXS). These techniques cannot solve the structure of the cement paste, but they can provide some characteristics of C-S-H:

2.5.4 Ca/Si ratio

Stoichiometry is the quantitative relationship between the reactants and products in a chemical reaction. The stoichiometry of C-S-H is not well defined and the dashed line between C, S, and H is to say that C-S-H compositions are known but the stoichiometry of C-S-H is not. To differentiate between different compositions, the ratio (Ca/Si) used can vary depending on cement composition (Taylor, 1997). In addition, the Ca/Si ratio varies irregularly from one point to another in the same cement paste sample, generally fluctuating between 1.5 and 2.3 with a mean of 1.75, but the value can be higher than 2.3 and lower than 1.2. Figure 11 shows the frequency of Ca/Si ratio over a 2.5 year duration.
Figure 11: Ca/Si ratio frequency histogram for C-S-H in Portland cement paste aged 1 day to 3 years (TEM microanalysis) (Richardson, 1999)

2.5.5 Water content

Water exists in hydrated cement paste in many forms. The best classification of different types of water is by Sierra (Gmira, 2003), which is based on structural and energy considerations. However, the dividing line between types is not always distinguishable. The different types are:

a) Interlayer water, the water between the layers of C-S-H, which is suggested to be a monolayer held by hydrogen bonding (Mehta and Monteiro, 2014). The C-S-H shrinks considerably when interlayer water is lost.

b) Adsorbed water is the water held on the surface of the hydrated pores by intermolecular forces, which decrease as the distance from the surface increases.

c) Capillary water is the water present in voids larger than 50Å (Mehta and Monteiro, 2014).

2.5.6 Layered structure

C-S-H has a layer structure that resembles Tobermorite and Jennite. The major difference between C-S-H and these minerals is the silicate chain. C-S-H has dimeric, pentameric and higher chains of SiO$_4$ in a sequence of 2, 5, 8, (3n-1). In the NMR study, SiO$_4$ represents one unit of the silicate chain, and these units can be connected in different forms of Q$^n$ where Q$^n$ represents the connectivity:
Q^0: isolated tetrahedral
Q^1: end tetrahedrals
Q^2: middle tetrahedral

During cement hydration, the Q^0 starts to decrease, accompanied by the formation of Q^1 and Q^2. The formation of Q^3 and Q^4 was not observed (Taylor, 1997). Figure 12 shows the different types of connectivity between the tetrahedral, from single tetrahedral (Q^1) to three units of tetrahedrals (Q^2).

![Different connections between tetrahedral SiO₄](image)

Figure 12: Different connections between tetrahedral SiO₄

During the first 24 hours of cement hydration, the Q^0 (monomer) starts to decrease, accompanied by the formation of Q^1 (dimer). At later ages, longer linear chains of species are formed; mainly pentamer and octamer following the 3n-1 rule. Dimer represents 40% of the total chain, even after six months of hydration. Figure 13 shows the distribution of ion species over time.
The mean silicate chain is the number of silicate tetrahedrals in a chain, calculated based on the ratio of $Q^2 / Q^1$

$$\text{Mean chain} = 2\left(1 + \frac{Q^2}{Q^1}\right) \quad (2.1)$$

Figure 14 shows the relationship between the mean silicate chain and Ca/Si. It clearly shows that the mean silicate chain decreases as Ca/Si ratio increases (Lagerblad et al., 1999).

![Figure 14: Mean silicate chain relationship with Ca/Si ratio (Hou, 2014)](image)

2.5.7 Mechanical properties

Nano-indentation is an experimental tool used to predict the mechanical properties of cement paste at nano- and micro-scale levels to show mechanical properties such as elastic modulus and hardness. Nano-indentation consists of pushing an indenter tip of a known geometry into the surface of any material. Load and penetration of the indenter are recorded and plotted on a graph, and indentation modulus (M) and hardness (H) are calculated. The applied load is in the micro-Newton range. Figure 14 shows that indentation modulus (M) is proportional to packing fraction. As packing increases along with indentation modulus (M), the agreement of the results between nano-indentation and
atomistic simulation increases as well. The relationship between Young modulus and the indentation modulus (M), bulk, shear modulus and Poisson ratio is provided by this equation:

\[ M = \frac{E}{1-\nu^2} = 4G \frac{3K+G}{3K+4G} \] (2.2)

Figure 15: Relationship between packing density and indentation modulus (Pellenq et al., 2009)

Young modulus is calculated from the indentation modulus, which is calculated from experiments. In regards to nano-indentation method, Canstantinides et al. (2013) found that there are two types of C-S-H: low density (LD) and high density (HD), with a packing factor of 0.64 and 0.74, respectively. The mean Young modulus of C-S-H is estimated to range between 20-30 GPa for LD and HD, respectively.

2.6 Mineral Analogy of C-S-H (ordered structure)

There are at least thirty crystalline structures that show similar composition to C-S-H. However, although the chemical compositions are similar, these minerals differ in their atomic arrangement, Ca/Si ratio, and the number of OH and H₂O. Most high-resolution TEM and extensive X-rays show that C-S-H has the structure of Tobermorite and Jennite (Taylor, 1997).
2.6.1 Tobermorite 14Å

Tobermorite 14Å (Ca₅Si₆O₂₆H₁₈) has SiO₄ tetrahedrals in a linear chain, twisted so they repeat at three tetrahedrals. This type of arrangement is called dreierkette. Tobermorite has a layer structure consisting of a central part of CaO₂ in which all the oxygen atoms are shared with the SiO₄ tetrahedrals. Only two tetrahedral share their oxygen with CaO₂; these are called paired tetrahedrals. The third is a bridging tetrahedral, which is positioned between the two paired tetrahedrals. The interlayer space between the individual layers is filled with Ca²⁺ and water molecules. Tobermorite is present in three different types, depending on water content: Tobermorite 14Å, 11Å and 9Å (Lea and Hewlett, 1998). Studies show that Tobermorite 14Å transforms into Tobermorite 11Å when heated to 80-100°C, and additional heating to 300°C transforms 11Å to 9Å (Lea and Hewlett, 1998). Tobermorite 11Å is present in two model structures: the Hamid structure has independent layers (Hamid, 1981), while the Merlino structure has chemically bonded layers (Merlino et al., 2001).

Figures 16, 17 and 18 show the structure of Tobermorite 14Å with details of pair and bridging tetrahedrals, interlayers filled with water and calcium, and the CaO₂ layer between the silicate chains. The X-ray diffraction of Tobermorite 14Å is shown in Figure 20.

![Idealized chemical structure of Tobermorite 14Å](image_url)

Figure 16: Idealized chemical structure of Tobermorite 14Å (Hewlett, 2004)
Figure 17: View of bridging tetrahedral and Pair tetrahedrals of Tobermorite (Shahsavari, Buehler, Pellenq, & Ulm, 2009)

Figure 18: Unit cell of Tobermorite 14A where Ca is presented in green, silicon in orange, oxygen in red, hydrogen in white, and the white line represents the unit cell
2.6.2 Tobermorite 11Å

Two types of Tobermorite 11Å crystal structure have been determined. Hamid (1981) and Merlino et al. (2001). According to Hamid, the Tobermorite structure in Figure 20 has a silicate chain in one layer independent of the other. However, in the Merlino structure presented in Figure 21, the silicate chain is connected and attached to the Ca-O layer. The Hamid structure comes in three forms that depend on the C/S ratio:

- C/S = 0.667 \(\text{Ca}_4[\text{Si}_6\text{O}_{14}(\text{OH})_4]\cdot2\text{H}_2\text{O}\)
- C/S = 0.830 \(\text{Ca}_5[\text{Si}_6\text{O}_{16}(\text{OH})_4]\cdot2\text{H}_2\text{O}\)
- C/S = 1 \(\text{Ca}_6[\text{Si}_6\text{O}_{18}]\cdot2\text{H}_2\text{O}\)

Figure 19: X-ray diffraction of Tobermorite 14Å

Figure 20: Hamid crystal structure showing linear silicate
2.6.3 Jennite

Jennite Ca$_9$(Si$_6$O$_{18}$H$_2$)(OH)$_6$·8H$_2$O is a natural mineral with a linear layer structure composed of SiO$_4$ tetrahedrals that are twisted so they repeat at intervals of three tetrahedrals. This arrangement is called dreierkette. The Ca/Si ratio of Jennite is 1.5 times higher than Tobermorite, and the layer has a thickness of 1.05 nm. The structure of Jennite has many similarities to Tobermorite, with a difference in the presence of the OH group in the sublayer CaO-OH and the absence of SiOH at the terminal.

When Jennite is heated to 70–90°C, the water molecules are lost and the unit cell shrinks in C direction (Bonaccorsi et al., 2004). The unit cell of Jennite and its X-ray diffraction is shown in Figures 22 and 23, respectively.

Figure 21: Merlino crystal structure showing silicate tetrahedral connected at the interlayer

Figure 22: Unit cell of Jennite where Ca is presented in green, silicon in orange, oxygen in red, hydrogen in white, and the white line represents the unit cell
The purpose of producing C-S-H (I) is to synthesize Tobermorite 14A. C-S-H (I) is the reaction product between calcium hydroxide and silica or between sodium silicate and calcium salts (Lea and Hewlett, 1998). C-S-H (I) has a structure similar to Tobermorite, with some missing bridging silicate tetrahedrals that follow the 3n-1 chain rule. C-S-H (I) has a Ca/Si molar ratio between 0.8-1.5 and a reduction of the mean silicate chain. On the other hand, C-S-H (II) is produced by hydrating C₃S or β-C₂S in excess water and repeatedly removing the liquid and replacing it with pure water until the Ca/Si ratio reaches the value of 2. Using electron diffraction, the C-S-H (II) shows a similar structure to Jennite with some disorder and finite chain length (Taylor, 1997).

2.7 Model of C-S-H at nano scale

To predict the performance and properties of concrete, it is important to have a model that explains the behavior of C-S-H under different conditions. The best-known models are discussed below:

2.7.1 Power and Brunauer (1964)

In the Power model, C-S-H is composed of particles held together by Van der Waals forces. Although the van der Waals interactions are weak, they can provide high concrete strength due to high surface
area (200 m$^2$/g). In addition, the model shows a fraction of chemical bonding (cross link) between the gel particles. The water in C-S-H can be categorized into three types:

a) Water combined chemically in the hydration product, which can only be evaporated at 1000°C.

b) Adsorbed water, which is the water held on the surface of the gel particles by physical forces (interlayer water).

c) The capillary water, which is free water inside the pores (Popovics, 1998).

The swelling in the Power model is caused by the presence of colloidal Ettringite, and the creep is caused by water squeezing between the particles, as shown in Figure 24.

![Figure 24: Schematic representation of Powers model (Feldman and Sereda, 1968)](image)

2.7.2 Feldman-Sereda model

The Feldman-Sereda model considers cement paste to be poorly crystalline layers of silicate, with the water having a more complex role than in the Power-Brunauer model. The water in the interlayer has a structural role, which is to connect two layers together and contribute to the rigidity of the system. The interlayer water can enter or leave the system (Feldman and Sereda, 1968). Figure 25 represents the Feldman-Sereda shematic model (Feldman and Sereda, 1968).
Figure 25: Schematic representation of Feldman-Sereda model (Feldman and Sereda, 1968)

2.7.3 Jennings model

Jennings proposed a colloidal model to explain the discrepancy between the results of investigations of the surface area and pore size distribution. The discrepancy does not only occur from one sample to another; results are different in the same sample. The model shown in Figure 27 is composed of basic blocks of 2.2 nm with equated shape; however, for simplicity, the block is shown as spherical. These blocks cluster to form units called globules, which are 5.6 nm in size. Jennings proposed two types of C-S-H: HD (High Density) C-S-H, where the structure is very compacted and N₂ (adsorption measurement) cannot penetrate, and LD C-S-H (Low Density), with loosely packed globules that are accessible by N₂. These two types of C-S-H have a gel porosity of 37% and 24% for LD and HD respectively, which can help explain the difference in surface area (Jennings, 2000). In addition, the pore sizes within these globules are similar for HD and LD and are not dependent on the volume of HD and LD from one cement to another (Jennings, 2000).
Figure 26: a) Basic building block, b) Globules, c) LD C-S-H, d) HD C-S-H and accessibility of N\textsubscript{2} to pores (Jennings, 2000)

2.7.4 Atomic model

In 1956, Taylor determined a relationship between the structures of C-S-H and Tobermorite made up of a central part with calcium hydroxide with missing hydrogen atoms. This sheet of CaO\textsubscript{2} is located between two parallel rows of silicate chains. These chains are twisted in a plan perpendicular to the CaO\textsubscript{2} sheet. Two thirds of the tetrahedrals are connected to the central chain of CaO\textsubscript{2} sharing oxygen atoms. The remaining third not connected to the central sheet is called a “bridging” tetrahedral. In
addition, between these layers, there is calcium and water molecule called the interlayer. Tobermorite 
$\text{Ca}(\text{Si}_6\text{O}_{18}\text{H}_2)\text{Ca}_4\text{H}_2\text{O}$ has a C/S ratio of 0.83 (5/6); in contrast C-S-H has an average C/S of 1.7. To 
increase the C/S ratio, Taylor proposed omitting the bridging tetrahedrals not directly connected to 
the central sheet CaO$_2$. Because the oxygen atoms are shared with the adjacent tetrahedral, the removal 
of the bridge leads only to the removal of SiO$_2$ and the transfer of the hydrogen atom to other position 
in the structure. In addition, the removed hydrogen atoms can be replaced by Ca atoms in the 
interlayer, as shown in Figure 27.

![Silicate chain of Tobermorite showing the probable position of H atoms and omission of bridging tetrahedrals](Taylor, 1986)

Thirty years later, Taylor proposed a C-S-H model that is a mixture of Tobermorite 14Å and Jennite 
structures to explain the high value of C/S ratio in the cement paste (1.7 to 2) as dimeric 1.4 nm 
Tobermorite and Jennite, with C/S ratios of 1.25 and 2.25 respectively. In both Tobermorite and 
Jennite, the silicate chain is considered infinite and the extreme C/S ratio of Jennite 
$\text{Ca}_9(\text{Si}_6\text{O}_{18}\text{H}_2)(\text{OH})_6\text{H}_2\text{O}$ is 1.5 (9/6) and 2.25 for dimer. Tobermorite has a C/S of 0.83 for infinite 
chain and 1.25 for dimer (Taylor, 1986). Figure 28 shows the relationship between Ca/Si ratio and 
silicate chain for Tobermorite and Jennite. At a chain length of two, the Ca/Si ratio is 1.25 and 2.25, 
respectively, and when the chain length is infinite, the ratio is 0.8 and 1.5, respectively. The C-S-H 
has a mean Ca/Si ratio of 1.75, which is between the value of dimer of Tobermorite and dimer of 
Jennite.
The mean silicate chain in C-S-H increases with time, and dimer are formed during the early age of hydration, while pentamer form at later ages. At early age of cement hydration, C-S-H can be similar to Torbermorite and Jennite-like structures, with dimer chain and C/S ratios of 1.25 to 2.25, respectively, as shown in Figure 28. This analogy has become the base of all other proposed models. Some researchers proposed another model based on the Taylor model. It is called Tobermorite/Portlandite, and in it, Tobermorite is sandwiched between Portlandite (Richardson, 2004) to increase C/S ratio. Other authors have proposed Tobermorite with more Ca$^{2+}$ and OH$^-$. Richardson (2004) proposed a generalized model which includes the Tobermorite-Jennite (T/J) and (T/CH) models. It also accounts for the substitution of Al$^{3+}$ for Si$^{4+}$ ions at the bridging tetrahedral. The formulation of this model is provided by the following formula:

\[
\{\text{Ca}_{2n}\text{H}_w\text{Si}_{(3n-1)}\text{O}_{(9n-2)}\}(\text{OH})_{w+n(y-2)}\text{Ca}_{(n,y)/2} \cdot m\text{H}_2\text{O}
\]

w: number of silanol; w/n: degree of protonation of silicate chain.

2.8 Other hydration products

2.8.1 Portlandite
Calcium hydroxide, also called Portlandite, forms from the hydration of C$_3$S and C$_2$S, consist of 20-25% of the volume of the hydration paste. Portlandite has a crystalline structure, in contrast to C-S-H, which has an amorphous one. The morphology of Portlandite (CH) is affected by space available.
for growth, temperature of hydration, and the presence of impurities and admixtures (Taylor, 1997).

In general, Portlandite (CH) has a hexagonal plate shape. In terms of mechanical contribution, it helps reduce pores in the paste, which has a small effect on paste strength. In addition, CH restrains the C-S-H from shrinkage. One characteristic of Portlandite is its solubility, which can cause leaching of CH when it contacts with water and can give aggressive agents access to the paste. On the other hand, it has the ability to protect concrete reinforcement from corrosion by increasing alkalinity (PH) in the system and creating a passive layer. The structure of Portlandite and its X-ray diffraction are shown in Figures 29 and 30, respectively.

![Figure 29: Unit cell Portlandite where Ca is presented in green, oxygen in red, hydrogen in white and the white line represents the unit cell](image1)

![Figure 30: X-ray diffraction of Portlandite](image2)
2.8.2 Ettringite

Ettringite [Ca$_3$Al(OH)$_6$12H$_2$O]$\cdot$(SO$_4$)$_3$2H$_2$O$]_2$ is the hydration product of Portland cement formed in limited amounts during early hydration. Ettringite contributes to the mechanical properties and stability of cement paste. Excessive amounts of Ettringite can cause expansion of the paste, which leads to cracking. The morphology of Ettringite depends on the space available and the supply of ions for growth. It can have a needle shape with a high ratio of length over diameter. Ettringite is formed from the reaction of C$_3$A with gypsum:

$$C_3A + 3C\bar{S}H_2 + 26H \rightarrow C_6AS_3H_{32}$$

Tricalcium Aluminate    Gypsum    water    Ettringite

The unit cell and X-ray diffraction of Ettringite are shown in Figures 31 and 32, respectively.

![Ettringite Unit Cell](image)

Figure 31: Unit cell Ettringite where Ca is presented in green, oxygen in red, hydrogen in white, aluminum in magenta and the white line represents the unit cell
Figure 32: X-ray diffraction of Ettringite

2.9 Summary

This chapter presents an overview of the clinker phases, hydration products of cement paste, similarities between C-S-H and Tobermorite and jennite, and models at nano-scale.

The major clinker phases are Alite (C₃S), Belite (C₂S), Tricalcium aluminate (C₃A) and Ferrite C₄AF. C₃S is the principal component of Portland cement clinker. It constitutes 50 to 90% of the clinker phases and is the most reactive composite. C₂S is the second principal clinker phase, and it makes up 10 to 40% of its composition. C₃A and C₄AF are the minor components in the clinker phases.

The chemical reaction between cement and water causes the setting and hardening of cement paste, a process called a hydration reaction. The hydration reaction is composed of several reactions that occur instantaneously or continuously. The degree of hydration is affected by a series of parameters such as the composition of the clinker phase, presence of foreign ions, surface area, w/c ratio, presence of SCM, chemical admixture and curing temperature. The hydration of C₃S and C₂S at normal temperatures leads to the formation of calcium silicate hydrate gel C-S-H and Portlandite (CH). The hydration of C₃S controls setting and hardening at early ages, while the hydration of C₂S is slow, with full hydration taking several years. The hydration of C₃A includes reaction with sulfate ions (gypsum), resulting in the formation of Ettringite. If the gypsum is consumed before the complete
hydration of C\textsubscript{3}A, the Ettringite reacts with the leftover C\textsubscript{3}A to form monosulfoaluminate. The hydration of C\textsubscript{4}AF is similar to C\textsubscript{3}A.

Concrete, mortar and cement paste are heterogeneous materials that present a multi-scale level in which the properties change as the length scale changes. Concrete, mortar and cement paste can be divided into three levels: macro-scale, micro-scale and nano-scale. At the macro-scale level, the aggregate and cement paste are the major phases, along with macro pores (entrapped air) if they are formed. At the micro- and nano-scale levels, each phase of the macro level shows heterogeneity. For instance, the calcium silicate hydrate appears to be dense in some areas (high density) (HD) and shows low density (LD) in others. In addition to C-S-H phases, Portlandite (CH) also can be detected in the capillary voids.

Although the structure of C-S-H is amorphous and is not yet solved, some techniques such as nuclear magnetic resonance (NMR), scanning electron microscope (SEM), small-angle X-ray scattering (SAXS) and nano-indentation tools assist researchers in determining some C-S-H characteristics. One of the most important characteristic of C-S-H is the molar ratio of CaO to SiO\textsubscript{2} (C/S), which is in the range of 1.2 to 2.3 with a mean of 1.75. With the nano-indentation tool, researchers can determine the mechanical properties of the two intrinsic types of C-S-H, which are LD C-S-H and HD C-S-H (low density and high density). The mean Young modulus of LD C-S-H and HD C-S-H are estimated to be between 20GPa and 30GPa. In addition, LD C-S-H and HD C-S-H show a packing factor of 0.64 and 0.74 respectively.

C-S-H has a layer structure in which the silicate chain follows the 3n-1 order. As hydration progresses, monomer numbers decrease and the dimer numbers increas. Higher chains such as pentamar and octamer are formed at late ages. Based on previous C-S-H modeling, a relationship has been established between C/S ratio and the mean silicate chain, in which the mean silicate chain increases when the C/S ratio decreases.

Tobermorite and Jennite are the structures that show the greatest similarities with C-S-H. Tobermorite can be found in three forms: Tobermorite 14A, 11A and 9A. Tobermorite 14A transforms to
Tobermorite 11A and 9A upon heating. Its structure is composed of SiO$_4$ tetrahedrons in a linear chain in a dreierkette form that repeats at every third tetrahedral, and a central layer of CaO$_2$ in which oxygen is shared with the SiO$_4$ and the interlayer is filled with water and calcium. Jennite also has a structure similar to Tobermorite, with the presence of the OH group in the sublayer CaO-OH and the absence of SiOH at the terminal. The C/S ratios of Tobermorite 14A and Jennite are equal to 0.83 and 1.5, respectively.
Chapter 3  Computational Methods

3.1 Introduction

The purpose of molecular modeling is to mimic the behavior of molecules with the use of computers to predict their behavior and properties at the molecular level. It has a wide range of applications for different disciplines such as material science, chemical engineering, and engineering science. Atomic simulation is the result of the solution of a mathematical model that estimates the position and response of electrons and nuclei. This model is based on two approaches: quantum and classic mechanics. The quantum approach is based on the solution of Schrödinger’s equation, where atoms and molecules are modeled through a mathematical equation. Schrödinger’s equation can be solved in two ways: the ab-initio method, where the properties of the structure are determined only by the solution of Schrödinger’s equation with the use of quantum mechanics, and by a semi-empirical method used to simplify the solution of Schrödinger’s equation using empirical parameters. Ab-initio method considers the electronic state and as result computation become expensive and rigorous. The semi-empirical method uses experimental results as an input to the mathematical model, with less computation time required than in the ab-initio method for a medium size system of 100 of atoms. The classic approach is based on classic physics and relies on the force field in which the parameters are embedded. It is a fast, useful method for large scale systems (with thousands of atoms), and does not deal with electronic properties. Also, it doesn’t require experimental data from ab-initio for the parameters.
3.2 Molecular mechanics

The molecular mechanics approach differs from the quantum mechanics approach in that the electrons and nuclei of the atoms are not explicitly included in the calculation. Molecular mechanics assumes molecules are a collection of masses that are connected to each other via springs of varying lengths (bond). These bond interactions can stretch, bend and twist. In addition, the non-bonded atoms interact with each other through Van der Waals and electrostatic interaction. The potential energy of the ensemble of atoms is calculated using Hooke’s law. As a result of these simplifications, molecular mechanics is a fairly fast computational method, and is practical for small and large molecules alike. Molecular mechanics is based on the Born-Oppenheimer approximations, which state that the Schrödinger equation can be divided into two equations of motion: one that describes the motion of the electrons, and another that describes the motion of the nuclei. These motions can be studied separately (Leach, 2001). Molecular mechanics is used to calculate the energy surface of molecules as a function of the motion of the nuclei. In molecular mechanics, the bonds have an “equilibrium” of lengths and angles. Force field parameters are the value of bond and angle, and the corresponding force constants are used in the potential energy functions. Any deviation from the equilibrium results in an increase of total energy of the molecule. The total energy of the system is the summation of the individual energy given by:

\[
\text{Energy} = \text{Stretching Energy} + \text{Bending Energy} + \text{Torsion Energy} + \text{Out of plane bending Energy} + \text{Non-Bonded Interaction Energy}
\]

3.3 Potential functions

3.3.1 Stretching energy

For simplicity, a harmonic oscillator was used to create a small vibration to calculate energy due to bond stretching from the equilibrium. A more accurate potential with higher terms can be used using the Taylor series. Morse potential is the most accurate for energy potential calculation. However, it has three parameters to calculate potential energy, which increases the computational cost for a
system with a large number of atoms. In addition, most of the stretch bond does not generally deviate significantly from its equilibrium. As a consequence, a harmonic oscillator that uses Hooke’s law is the best approach for energy calculation. Figure 33 shows bond stretching and Hook’s law equation to calculate energy.

\[ E_{\text{stretch}} = \frac{K_b}{2} (r - r_0)^2 \quad (3.1) \]

Figure 33: Bond stretching

Where \( K_b \) is the stiffness of the bond spring, \( r_0 \) is the equilibrium bond length and \( r \) is the current bond length after vibration.

3.3.2 Bending energy

Figure 34 shows the energy due to bending of the bond between three atoms joined in a series caused by the deviation of the angle from their reference. It can be calculated as follows:

\[ E_{\text{bending}} = \frac{K_\theta}{2} (\theta - \theta_0)^2 \quad (3.2) \]

Figure 34: Angle bending

\( K_\theta \) is the force constant, \( \theta_0 \) : Angle at equilibrium , \( \theta \) : Instantaneous angle

Less energy is required to distort the angle between the bonds than to stretch or compress a bond. A more accurate potential energy can be calculated using the Taylor series; it incorporates high order terms, which leads to a higher computational cost.
3.3.3 Torsion energy

Most energy deviation is due to torsional and non-bonded energy contributions. Torsion energy is the result of deviation of the angle for a set of four atoms. Torsion energy is modelled by a simple periodic function, as shown in Figure 35.

\[
E_{\text{Torsion}} = \sum_{n=1}^{\infty} \frac{E}{2} (\cos n\phi - \Theta) \tag{3.3}
\]

Figure 35: Bond torsion

Where \( E \): barrier height, \( n \): multiplicity, \( \phi \) torsion angle and \( \Theta \): phase shift

3.3.4 Out of plane bending

Deviation of the atom out of plane is treated differently than just bending on the plane. In out of plane, the three bending atoms are in one plane and the fourth atom is bent, making an angle \( \omega \) with the plane. The energy deviation can be presented as shown in Figure 36 and expressed as harmonic function as follows:

\[
E_{\text{out plane}} = \frac{K_\Theta}{2} (\omega)^2 \tag{3.4}
\]

Figure 36: Out of plane

Where \( K_\Theta \) is the force constant, \( \omega \) is the angular deviation from the plane.

3.3.5 Non-bonded interactions

Atoms that are not bonded to each other interact through non-bonded interaction forces. These interactions occur in space and are dependent on the distance between particles. Non-bonded
interactions can be divided into two categories: electrostatic interactions and van der Waals interactions.

\[ E_{\text{non-bonded}} = E_{\text{electrostatic}} + E_{\text{Van der waals}} \]  

(3.5)

3.3.6 Electrostatic interactions (long range)

Due to the difference in electronegativity between atoms, unequal distribution of charge is created. These charges produce an electrostatic force between atoms. The electrostatic interaction between two atoms \( i \) and \( j \) with partial charge is calculated using Coulomb’s law.

\[ E_{\text{electrostatic}} = \sum_{j=1}^{Na} \sum_{i=1}^{Nb} \frac{q_i q_j}{4\pi \epsilon r_{ij}} \]  

(3.6)

Where \( q_i \) and \( q_j \) are the charges

\( r_{ij} \) is the distance of the separation

\( \epsilon \) is the permittivity

\( Na, Nb \): number of punctual charges

Electrostatic interaction (Figure 37) has a slower rate of decay \((1/r)\) than Van der Waals, which decays at a rate of \(1/r^6\). It requires a considerable distance for decay, which leads to high computational cost.

![Figure 37: Electrostatic interactions](image-url)
3.3.7 Van der Waals interactions (short Range)

In addition to electrostatic interaction, Van der Waals interactions occur due to the induced dipole between two or more molecules. Van der Waals are the weakest interactions, however; due to the presence of many van der Waals interactions between atoms or molecules, they become very strong. The Van der Waals interaction has a short-range effect. As atoms move away from each other, energy decays rapidly at a rate of $1/r^6$. The van der Waals interaction energy is composed of two terms: attractive and repulsive. Repulsion occurs when the distance between two attracting particles becomes very small, which causes a high increase in energy due to an overlap of electrons in the space (Pauli). The attractive force occurs due to the instantaneous dipole from the fluctuation of electron clouds. The mathematical model that best describes this attraction and repulsion is called the Leonard Jones Potential [L-J] or 6-12, as shown in Figure 38 (Leach, 2001).

$$E_{vdw} = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{12} \right]$$  (3.7)

![Figure 38: The Lennard-Jones potential (Leach, 2001)](image)

3.3.8 Force field used

Force field is the functional form and parameter set used to calculate the potential energy of a system of atoms in molecular mechanics and molecular dynamics simulations. It is the core of material characterization using molecular mechanics. Different force field types such as UNIVERSAL,
DREIDING and CVFF have been used in literature to simulate minerals such as clinker phases and hydration products. However, COMPASS, which is the most recent force field added to the Materials Studio software, has more reliable results compared to other force fields.

3.3.9 COMPASS force field

The force field used in molecular mechanics is the major parameter affecting the accuracy of the results. Among the different force fields used, COMPASS (Condensed-phase Optimized Molecular Potentials for Atomistic Simulation Studies), is the most recent force field added to the material software. It is a mixture of the ab-initio method and classic mechanics, which is the reason for its special characteristics and accurate predictions. It has been broadly used in a wide range of molecules and polymers.

\[ E_{total}^{\text{compass}} = E_{valence} + E_{cross term} + E_{non-bond} \]

\[ = \sum \{ K_2 (b - b_0)^2 + K_3 (b - b_0)^3 + K_4 (b - b_0)^4 \} \]

\[ + \sum_\theta \{ H_2 (\theta - \theta_0)^2 + H_3 (\theta - \theta_0)^3 + H_4 (\theta - \theta_0)^4 \} \]

\[ + \sum_\Omega \{ V_1 [1 - \cos(\Phi - \Phi_0)] + V_2 [1 - \cos(2\Phi - \Phi_0)] + V_3 [1 - \cos(3\Phi - \Phi_0)] \} \]

\[ + \sum_z K_z X^2 + \sum_b \sum_{b'} F_{bb'} (b - b_0)(b' - b'_0) + \sum_\theta \sum_{\theta'} F_{\theta\theta'} (\theta - \theta_0)(\theta' - \theta'_0) \]

\[ + \sum_b \sum_{\theta} F_{b\theta} (b - b_0)(\theta - \theta_0) + \sum_b \sum_\Phi F_{b\Phi} (b - b_0) \{ V_1 \cos \Phi + V_2 \cos 2\Phi + V_3 \cos 3\Phi \} \]

\[ + \sum_{b'} \sum_\Phi F_{b'\Phi} (b' - b'_0) \{ V_1 \cos \Phi + V_2 \cos 2\Phi + V_3 \cos 3\Phi \} \]

\[ + \sum_\theta \sum_\Phi F_{b\Phi} (\theta - \theta_0) \{ V_1 \cos \Phi + V_2 \cos 2\Phi + V_3 \cos 3\Phi \} \]

\[ + \sum_\Phi \sum_\Omega \sum_{\theta'} k_{\theta\theta'\Phi} \cos \Phi (\theta - \theta_0)(\theta' - \theta'_0) + \sum_{i>j} \frac{q_i q_j}{r_{ij}} + \sum_{i>j} \frac{A_{ij}}{r_{ij}^2} \]

Where: K, H, F and B are constant, b is bond distance, \( \theta \) is the bond angle, \( \Omega \) torsion angle and \( V \) is coefficient of coefficient of cosine Fourier expansion (Materials Studio software V7).
3.4 Energy Minimization

Geometry optimization is the first important step in molecular computation. The purpose of a geometry optimization is to find a minimum energy potential surface with respect to atomic position. Any movement of the structure far from its equilibrium will generate a configuration with higher energy. The geometry of the molecule determines many of its physical and chemical properties (Ramachandran et al., 2008). The potential energy function can have many minimum values. The lowest is called the global energy minimum. To locate these energy minimum points, minimization algorithms are used. They can be categorized into two groups: one that uses a derivative of the energy function with respect to coordinates and one which does not. The direction of the derivative (positive or negative) indicates where the minimum energy is located, and the magnitude of the derivative indicates steepness. The energy of the system can be lowered by moving atoms to the relaxed position. The best algorithm is one that finds the minimum location as quickly as possible. Most algorithms move downhill of the surface energy, so they only locate the minimums that are nearest, as shown in Figure 39 and illustrated by the following function (Leach, 2001):

\[
F = -\frac{\partial E}{\partial r}
\]

(E: potential energy, \(r\): atom position)

Figure 39: A schematic one-dimensional energy surface, minimization method moving downhill to nearest minimum (Leach, 2001)
3.4.1 Steepest Descent (SD)

The Steepest Descent (SD) minimization algorithm uses the first derivative to find the position of atoms with minimum energy. The force derived shows the direction in which the atoms have to move towards the relaxed structure. The energy potential is calculated from the initial geometry; the atoms move with small increments in the direction of the force (called line of search) until the minimum is found in that direction. Another line search, perpendicular to the previous minimum point, locates the lowest minimum, as shown in Figure 40. The coordinates at step $i$ are:

$$x_{i+1} = x_i + \lambda x_i \quad (3.9)$$

![Figure 40: Minimization path followed by steepest-descents approach for simple quadratic function (Materials Studio)](image)

This algorithm is effective when the initial configuration is far from the minimum. However, it will not be effective if the gradient of the potential is small.

3.4.2 Conjugate gradient

The Steepest Descent method converges slowly to the minimum because each line search segment must be orthogonal to the previous one, and therefore the progress made by the first line search is lost. In the conjugate gradient method, the direction mutually conjugates such successive steps to continue refining the direction towards the minimum. It uses both the current gradient and the previous search direction to drive minimization. Because the conjugated gradient method uses the minimization history to calculate search direction and contains a scaling factor for determining step
size, this method converges faster and makes step sizes optimal as compared to the Steepest Descent technique. Figure 41 shows the minimization comparison between the steepest descent and conjugate gradient methods.

![Minimization comparison between Steepest Descent and Conjugate gradient method](image)

Figure 41: Minimization comparison between Steepest Descent and Conjugate gradient method (Materials Studio)

### 3.5 Molecular dynamic

Molecular dynamics is a computer simulation used to simulate the movement of atoms and molecules as they interact with each other (Frenkel & Smit, 2002). The MD simulation produces a trajectory of the atoms and molecules that provides their positions and velocities in the system as a function of time by numerically solving Newton’s law equation:

$$\frac{d^2 x_i}{dt^2} = \frac{F_{xi}}{m_i}$$  \hspace{1cm} (3.10)

Where $F_{xi}$ is the force acting on the particles due to the interactions, $m_i$ mass of the particles and $x_i$ coordinates of atoms.

The purpose of molecular dynamics is to calculate the force between particles and their energy. MD is also used to determine the average properties of the system at a macroscopic level, which can be
compared with experiments that have been conducted and also those that it is not possible to perform (Leach, 2001).

At a specified temperature, an initial random velocity is applied to the atoms, which causes the atoms to change position (trajectory). With potential (force field) used, the forces acting on the atoms can be determined at their new positions by:

\[ F_i = \frac{dE(r_i)}{dr_i(t)} = \nabla E(r_i) \quad (3.11) \]

Where \( E \) is energy potential calculated from classic mechanics, \( r_i \) is the position of atoms. When velocity is applied to the set of molecules, the system will be out of equilibrium (thermodynamic equilibrium), and with time will be relaxed to a new value. When the system is brought to equilibrium, measurements of system properties can then be calculated. The force calculation is completed in a series of short steps (1 fs), and in each step the force, the new position and velocities are calculated until equilibrium is reached, the trajectory is obtained and average properties are calculated:

\[ <A> = \frac{1}{M} \sum_{i=1}^{M} A\left(P^N, r^N\right) \quad (3.12) \]

Where \( M \) is number of time steps, \( P \) is momentum of atoms, \( N \) number of atoms, \( r \) is the position of the atom.
The flow chart of the MD simulation algorithm is shown in Figure 42.

3.5.1 Integration of equation of motion (algorithms)

The force on each particle will change as the particle changes its position, or the particles interacting with it change position. The motions of all particles are connected to each other, which creates a multi-body equation that cannot be solved analytically. Instead, the equation of motion is solved by using the finite difference method. The main idea is to break the integration into small steps at $\Delta t$. 
The Verlet algorithm works in two steps: 1) use the current position of the particles to calculate the current force applied to the particles 2) use the current position \( r_n \) and previous \( r_{n-1} \) with the current force calculated in step 1 to calculate the position in the next step \( r_{n+1} \). The algorithm is based on the third expansion of the Taylor series for forward and backward expansion, which calculates the position at \( t + \Delta t \) (Becker et al., 2001).

\[
\begin{align*}
  r(t + \Delta t) &= r(t) + \Delta t v(t) + \frac{1}{2} \Delta t^2 a(t) + \cdots \quad (3.13) \\
  r(t - \Delta t) &= r(t) - \Delta t v(t) + \frac{1}{2} \Delta t^2 a(t) + \cdots \quad (3.14)
\end{align*}
\]

\[ (2.13) + (2.14) \quad r(t + \Delta t) = 2r(t) - r(t - \Delta t) + \Delta t^2 a(t) \quad (3.15) \]

The velocity can be calculated as:

\[ (2.13) - (2.14) \quad v(t) = \frac{[r(t + \Delta t) - r(t - \Delta t)]}{2\Delta t} \quad (3.16) \]

Figure 43 is a graphical representation of the steps followed by the Verlet algorithm:

![Graphical representation of steps followed by Verlet algorithm](image)

Figure 43: Graphical representation of steps followed by Verlet algorithm (Allen and Tildesley, 1987)

3.5.2 Periodic boundary condition (PBC)

When we simulate a system, the atoms at the face of the simulated system have fewer neighbors than the atoms inside the box; the applied force on molecules at the surface is different than on those in the bulk. As a result, the system will not reflect the reality, no matter how large it is. Periodic boundary condition is used to solve this issue; in it, a system is replicated as an imaginary box to infinity in all Cartesian directions. During the simulation, as particles move in the original box, the particles in the periodic image will move the same way as in the original box. In addition, if one particle passes through the face of the original box, it reappears on the opposite face of the imaginary box with the
same velocity, which leads to a constant number of particles in the box and maintained density. During the simulations, the particles are free to cross the six cube faces, as shown in Figure 44.

![Figure 44: Periodic boundary conditions in molecular dynamics (Allen and Tildesley, 1987)](image)

3.5.3 Cutoff distance

Due to the elimination of boundary conditions, the number of interactions between atoms increases, which leads to a reduced speed calculation. A cut-off distance is used to reduce the number of interactions, and any pair interaction beyond the cut-off distance is ignored. The cut-off distance should be less than half of the box dimension (Cf < L/2).

3.5.4 Ensembles

The purpose of molecular modelling is to determine the macroscopic properties of a system (pressure, energy, volume) through microscopic simulation. Statistical mechanics are able to determine the macroscopic properties from the individual atoms or molecules making up the system. At the macrocosmic level, a system can be defined by parameters such as temperature (T), pressure (P), Volume (V), and number of particles (N). At the microscopic level, a system is defined by the position of atoms (q) and momenta (p), which are considered the coordinates in a three-dimensional space.
called phase space. An ensemble is a collection of all possible systems which have different microscopic states but identical macroscopic ones. Different ensembles can be used in molecular dynamic simulation.

In statistical mechanics, the ensemble average is given by:

\[
\langle A \rangle = \int \int A(p, q)P(q, p)dpdq
\]

\(P(q,p)\) is the probability of the system to be at the phase space point \((q,p)\), and \(A(q,p)\) is the system property value at that point.

3.5.5 NVE

A constant number of particles, constant volume and constant energy ensemble (NVE). The Newton equation of motion is solved without any temperature or pressure control.

3.5.6 NVT

A constant number of particles, constant volume, and constant temperature ensemble (NVT) maintain a constant temperature where atom velocities have to be controlled.

3.5.7 NPT

A constant number of particles, constant pressure, and constant temperature ensemble (NPT) control both temperature and pressure. The unit cell is allowed to change and pressure is adjusted by adjusting the volume.

3.6 Mechanical Properties

The most important mechanical properties of cement-based material are the short-term deformations controlled by the elastic modulus. The elastic properties describe how the material responds to small reversible deformation. The relationship between stress and strain can be written using Hooke’s law:

\[
\sigma_i = \sum_{j=0}^6 C_{ij} E_j \quad (3.17)
\]

Where \(C_{ij}\) are elastic tensor coefficients, \(E_j\) is the strain.
In molecular modeling, the elastic constants \( (C_{ij}) \) can be calculated by the second derivative of energy density \( (\text{Energy/Volume}) \) with respect to lattice strain components.

\[
C_{ij} = \frac{1}{V} \left( \frac{\partial^2 U}{\partial \varepsilon_i \partial \varepsilon_j} \right)
\]  
(3.18)

Where: \( \varepsilon_i, \varepsilon_j \) are lattice strain components, \( U \) is potential energy, and \( V \) is simulation cell volume.

The elastic compliance matrix \( (S) \) can be obtained by finding the inverse matrix of \( C_{ij} \)

3.6.1 Bulk modulus and shear modulus

Bulk (K) and shear (G) modulus can be calculated from the elastic tensor coefficient. There are three ways to calculate K and G. The K and G in the Reuss are:

\[
K_{\text{Reuss}} = \left[ (S_{11} + S_{22} + S_{33}) + 2(S_{12} + S_{13} + S_{23}) \right]^{-1}
\]  
(2.19)

\[
G_{\text{Reuss}} = \frac{15}{4} \left[ (S_{11} + S_{22} + S_{33} - S_{12} - S_{13} - S_{23}) + 3(S_{44} + S_{55} + S_{66}) \right]^{-1}
\]  
(3.20)

The Voight definitions of K and G are:

\[
K_{\text{Voight}} = \frac{1}{9} \left[ (C_{11} + C_{22} + C_{33}) + 2(C_{12} + C_{13} + C_{23}) \right]
\]  
(3.21)

\[
G_{\text{Voight}} = \frac{1}{15} \left[ (C_{11} + C_{22} + C_{33}) + 3(C_{44} + C_{55} + C_{66}) - C_{12} - C_{13} - C_{23} \right]
\]  
(3.22)

The bulk (K) and shear (G) modulus from Reuss and Voight represent the upper and lower limits.

The average of the value of Reuss and Voight represents the Hill average

\[
K_{\text{Hill}} = \frac{K_{\text{Voight}} + K_{\text{Reuss}}}{2}
\]  
(3.23)

\[
G_{\text{Hill}} = \frac{G_{\text{Voight}} + G_{\text{Reuss}}}{2}
\]  
(3.24)
When bulk and shear have been calculated, Young modulus (E) and Poisson ratio are then calculated, assuming an isotropic material:

\[ E = \frac{9G}{3+G/K} \]  (3.25)

\[ \nu = \frac{3-2G/K}{6+2G/K} \]  (3.26)

3.7 Summary

This chapter describes different molecular computation methods and concepts used in literature. Two approaches are defined: molecular modeling based on quantum mechanics and molecular modeling based on molecular mechanics. One of the most famous methods that uses quantum mechanics is called ab-initio, which is based on the solution of the Schrödinger equation. Different techniques have been derived from this approach. However, because quantum mechanics consider the electronic state, the computation becomes expensive, rigorous and long. In the second approach, which uses classic (molecular) mechanics, the atoms are considered a collection of mass, connected with springs that permit the calculation of the potential energy of the system.

The force field used in molecular mechanics is a major parameter that can the reliability of results. COMPASS force field, the most recent force field used in cement paste simulation, has been previously used in different ranges of molecules and polymers simulations.

Molecular modeling with COMPASS force field is used to locate the points at which the energy of atoms is at its minimum; any movement from these points generates additional energy in the system. Energy minimization is used to locate the minimum energy of a system using algorithms such as Steepest Descent (SD), Conjugate Gradient and others. Energy minimization is also used as a pre-equilibrium before running molecular dynamics.
Molecular dynamics simulate the progress and interaction of atoms over a period of time. The Verlet algorithm is used to numerically solve the Newton equation of motion. Molecular dynamics act as a bridge between microscopic and macroscopic scales in which we can obtain bulk properties.

Elastic constants can be calculated by the numerical second derivative of energy density (Energy/Volume) with respect to lattice strain components.
Chapter 4  Elastic Properties of major clinker phases

4.1  Introduction

When cement is mixed with water, it results in a paste in which the different crystal phases are composed from reacted and unreacted cement. The reacted components, called hydration products, are mainly composed of calcium silicate hydrate (C-S-H) and Portlandite (CH), in which C-S-H is the main contributor to the mechanical properties of the cement paste. The unreacted components (C_3S, C_2S, C_3A, C_4AF) are composed mainly from the clinker phase and have high percentages of Alite and Belite. The anhydrate (unreacted) phase can be found in early ages of cement paste, or in mature ages when W/C ratio is low. The anhydrate phase in this case can also affect global mechanical properties. Knowing the properties of these different phases is important to improving paste quality. Calculating mechanical properties such as Young modulus, Poisson ratio, bulk modulus and shear modulus is important at the nano-scale level to determine properties at macro- and micro-scale levels.

4.2  State of art

Velez et al. (2001) performed experiments using the nano-indentation method at the microscopic level to determine the elastic modulus and hardness of synthetic C_3S, C_2S, C_3A and C_4AF. They also used resonance frequency measurement at the macroscopic level. The structure and purity of the synthesized phases were analyzed by X-ray diffraction. In the nano-indentation experiment, the penetration depth of the indenter was about 300 to 500 nm, with an applied load of 40 to 50 mN. They
found that the elastic modulus of different synthetic clinker phases is in the range of 125 to 145 GPa (Velez et al. (2001)). They also observed no significant change in Young modulus between C₃S and Alite which has some impurities. The same applied for C₂S and Belite. For resonance frequency testing, specimens with different densities were made to account for porosity. The elastic modulus (as a function of porosity) was calculated by measuring the resonance frequency of the bar specimen (10x1x2cm), and they found that the elastic modulus at zero porosity was higher, in a range of 140 to 160GPa, for all clinker phases. They also concluded that elastic modulus decreases about 30 to 40% when porosity increases from 10 to 20%. For C₃S, the fit equation for porosity is

\[ E = 146.89 * e^{-3.4p} \]

Mondal (2008) performed experiments to determine the variation of the elastic modulus of C-S-H throughout the surface. He used a triboindenter instrument, which combines nano-indentation and scanning probe microscopy (SPM) imaging. The advantage of this instrument is its ability to locate the indenter in a specific location with a high-resolution image, identify the different phases of the cement paste, and position the indenter probe within 10nm (Mondal 2008). Mondal (2008) found that Young modulus of the anhydrate cement paste is around 100GPa, which is slightly lower than the Velez results (Velez et al., 2001). They also found that there are three groups of elastic modulus that decrease with distance from the anhydrate cement. The elastic modulus map using nano-indentation is presented in Figure 45.

Figure 45: Elastic modulus map using nano-indentation (Mondal, 2008)
The elastic modulus map is based on the elastic modulus of each indentation. Four groups of elastic modulus were found: anhydrate cement, low stiffness C-S-H, high stiffness areas, and very low stiffness areas due to C-S-H porosity.

4.3 Basic review of crystal structure

A crystal is build up from a repeating structural motif (Atkins, 1986), which can be atoms or molecules. A unit cell is an imaginary parallelepiped that contains one unit of the repeating pattern. The length of the unit cell is a, b and c, and the angles between them are $\alpha$, $\beta$ and $\gamma$. A unit cell can be classified into seven crystal systems: triclinic, monoclinic, orthorhombic, rhombohedral, tetragonal, hexagonal and cubic. Table 1 shows the different types of crystals including cubic, monoclinic, triclinic and their axial parameters (length and angles) a, b, c, $\alpha$, $\beta$, and $\gamma$, as presented in Figure 46.

Table 1: Crystal system with lattice parameters

<table>
<thead>
<tr>
<th>Crystal family</th>
<th>Crystal system</th>
<th>Axial relationships</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isometric</td>
<td>Cubic</td>
<td>$a = b = c$; $\alpha = \beta = \gamma = 90^\circ$;</td>
</tr>
<tr>
<td>Tetragonal</td>
<td>Tetragonal</td>
<td>$a = b \neq c$; $\alpha = \beta = \gamma = 90^\circ$;</td>
</tr>
<tr>
<td>Orthorhombic</td>
<td>Orthorhombic</td>
<td>$a \neq b \neq c$; $\alpha = \beta = \gamma = 90^\circ$;</td>
</tr>
<tr>
<td>Monoclinic</td>
<td>Monoclinic</td>
<td>$a \neq b \neq c$, $\alpha = 90^\circ$, $\beta \neq 90^\circ$, $\gamma = 90^\circ$;</td>
</tr>
<tr>
<td>Anorthic</td>
<td>Triclinic</td>
<td>$a \neq b \neq c$, $\alpha \neq 90^\circ$, $\beta \neq 90^\circ$, $\gamma \neq 90^\circ$;</td>
</tr>
<tr>
<td>Hexagonal</td>
<td>Hexagonal</td>
<td>$a = b \neq c$, $\alpha = \beta = 90^\circ$, $\gamma = 120^\circ$;</td>
</tr>
<tr>
<td></td>
<td>Trigonal or</td>
<td>$a = b = c$, $\alpha = \beta = \gamma$; or</td>
</tr>
<tr>
<td></td>
<td>Rhombohedral</td>
<td>$d = b' \neq c'$, $\alpha' = \beta' = 90^\circ$, $\gamma' = 120^\circ$;</td>
</tr>
</tbody>
</table>

(hexagonal axes)

Figure 46: Crystal structure showing angles in the cell
4.4 Computation method

The goal in this study was to use COMPASS force field instead the other current force fields in the Materials Studio software (Materials Studio v7.0). Elastic properties of all the anhydrite phases of cement were calculated by using COMPASS force field as the first choice. However, COMPASS was not applicable for all the clinker phases; DREIDING and CVFF in the Materials Studio software was also used when COMPASS was not applicable. Before starting molecular dynamic simulation, it is a prerequisite to do an energy minimization of the structure to remove any excess energy in the system. Minimization (or optimization of the geometry) is an iterative process in which the coordinates of the atoms and cell parameters are adjusted and bring the structure to a minimal energy. Minimization leads to a structure that is similar to the real physical structure. In addition, when performing a geometry optimization, temperature and pressure are not considered. However, they are controlled in molecular dynamics (MD). Materials Studio software uses a smart minimizer algorithm, which starts with steepest descent and is followed by the conjugated gradient algorithm (Materials Studio v7.0).

After energy minimization of the structure, the dynamic simulation was performed in two steps: equilibrium and production. At the beginning of the equilibrium phase, a random velocity was assigned to all atoms in the structure at a selected target temperature. The temperature was controlled during the equilibrium phase by nose thermostat, and the pressure was controlled by Anderson barostat. Molecular dynamics can be completed in a different ensemble: NVE, NVT, NPH and NPT. All molecular dynamics simulations in this study were completed in the NPT ensemble, in which \( N \) is the number of atoms, \( V \) is the volume of periodic cell, \( E \) is the total energy is temperature, \( P \) is the pressure and \( H \) is enthalpy. The equilibration stage was used to equilibrate the kinetic and potential energies. The kinetic energy was increased due to the heating of the system from 0°C to 300°C, and had to be equilibrated by potential energy. As soon as kinetic energy is equal to the potential energy, equilibration is reached. Plotting energy/temperature vs. time determines whether the system has reached that stage. When quantities such as energy and temperature fluctuated around their average, the system reached its equilibrium, as shown in Figures 47 and 48.
Figure 47: Potential energy, kinetic energy, non-bonded energy and total energy fluctuation during equilibration phase

Figure 48: Fluctuation of temperature and running average temperature during equilibration phase

When the equilibrium of the system reached the desired temperature (298k) and pressure, the production run started and data and statistics were collected. Another important parameter in MD simulation is the time step used in algorithm integration; too large a step can lead to inaccurate results. NPT ensemble was used in all simulation in this study.

4.4.1 Crystal parameters

Table 2 shows the lattice parameters (a, b, c, α, β, and γ) of the clinker phases used in this study.
Table 2: Unit cell parameters for different clinker phases

<table>
<thead>
<tr>
<th>Crystal lattice</th>
<th>C3S</th>
<th>β-C2S</th>
<th>C3A</th>
<th>C4AF</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>11.6389</td>
<td>5.480</td>
<td>15.263</td>
<td>5.584</td>
</tr>
<tr>
<td>b</td>
<td>14.1716</td>
<td>6.760</td>
<td>15.263</td>
<td>14.600</td>
</tr>
<tr>
<td>c</td>
<td>13.6434</td>
<td>9.280</td>
<td>15.263</td>
<td>5.374</td>
</tr>
<tr>
<td>a</td>
<td>104.982</td>
<td>90.000</td>
<td>90.000</td>
<td>90.000</td>
</tr>
<tr>
<td>β</td>
<td>94.622</td>
<td>94.330</td>
<td>90.000</td>
<td>90.000</td>
</tr>
<tr>
<td>γ</td>
<td>90.107</td>
<td>90.000</td>
<td>90.000</td>
<td>90.000</td>
</tr>
<tr>
<td>Ref</td>
<td>[1]</td>
<td>[2]</td>
<td>[3]</td>
<td>[4]</td>
</tr>
</tbody>
</table>

Ref: reference
[1]: (Torre et al., 2008)
[2]: (Midgley, 1952)
[3]: (Mondal and Jeffery, 1975)
[4]: (Jupe et al., 2001)
[5]: (Pedersen and Semmingsen, 1982)

4.4.2 Geometry minimization

An empirical approach was used based on COMPASS force field, where applicable, to determine interactions between atoms. However, this targeted force field was not applicable for all clinker phases. After creating the cell with the atoms connected, a geometry optimization was run to bring the structure to its relaxed state. Because COMPASS force field showed limitations in the case of C3A and C4AF, DREIDING and CVFF force fields, respectively, were used for those two phases. Table 4 shows the parameters used to optimize the structure.

Table 3: Unit cell parameters for different clinker phases

<table>
<thead>
<tr>
<th>Geometry minimization parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Force field</td>
</tr>
<tr>
<td>Quality</td>
</tr>
<tr>
<td>Summation Method</td>
</tr>
<tr>
<td>Repulsive cutoff</td>
</tr>
</tbody>
</table>
4.4.3 Dynamic simulation parameters

After minimizing the structure, a dynamic simulation was run where the atoms interacted with each other for a period of time. The position of the atoms was determined by numerically solving the Newton equation, and the forces between atoms were calculated with the potential energy using the force field mentioned in Table 4. Table 5 shows the parameters used in dynamic simulations.

<table>
<thead>
<tr>
<th>Dynamic Simulation parameters</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ensemble</strong></td>
<td>NPT</td>
</tr>
<tr>
<td><strong>Temperature</strong></td>
<td>298 K (25°C)</td>
</tr>
<tr>
<td><strong>Pressure (GPa)</strong></td>
<td>0.0001</td>
</tr>
<tr>
<td><strong>Time step (fs)</strong></td>
<td>0.5</td>
</tr>
<tr>
<td><strong>Total simulation (ps)</strong></td>
<td>200-500</td>
</tr>
<tr>
<td><strong>Thermostat</strong></td>
<td>Nose</td>
</tr>
<tr>
<td><strong>Barostat</strong></td>
<td>Anderson, Berendsen</td>
</tr>
</tbody>
</table>

4.5 Computation results

When the forces between atoms are calculated, the mechanical properties of the structure can be determined. Table 5 shows the mechanical properties of the clinker phases from atomic simulation compared with other theoretical and experimental results from literature.
Table 5: Elastic properties of cement clinker: bulk modulus, shear modulus and Young modulus, Poisson ratio and experimental results (Appendix I, II, III and IV)

<table>
<thead>
<tr>
<th>Crystal</th>
<th>C₃S</th>
<th>C₂S</th>
<th>C₃A</th>
<th>C₄AF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Modulus: K (GPa)</td>
<td>162.3</td>
<td>64.6</td>
<td>176.1</td>
<td>127.8</td>
</tr>
<tr>
<td>Shear modulus: G (GPa)</td>
<td>50.04</td>
<td>50.6</td>
<td>64.2</td>
<td>82.9</td>
</tr>
<tr>
<td>Young modulus: E (GPa)</td>
<td>136.1</td>
<td>120</td>
<td>171.7</td>
<td>204.5</td>
</tr>
<tr>
<td>ν: Poisson ratio</td>
<td>0.36</td>
<td>0.19</td>
<td>0.34</td>
<td>0.234</td>
</tr>
<tr>
<td>Forcefield used</td>
<td>FC</td>
<td>FC</td>
<td>FD</td>
<td>FCV</td>
</tr>
<tr>
<td>Ref results (GPa)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K[8]</td>
<td>103</td>
<td>111</td>
<td>133.26</td>
<td>175.6</td>
</tr>
<tr>
<td>G[8]</td>
<td>54.5</td>
<td>53.1</td>
<td>63.9</td>
<td>93.5</td>
</tr>
<tr>
<td>E[8]</td>
<td>138.9</td>
<td>137.9</td>
<td>165.1</td>
<td>238.2</td>
</tr>
</tbody>
</table>

[6]: (Velez et al., 2001)  
[7]: (Bass, 1995)  
[8]: (Manzano et al., 2009)  
FC: Force field COMPASS  
FD: Force field DREIDING  
FCV: Force field CVFF

The atomic simulations of C₃S, C₂S, C₃A, C₄AF structures were conducted using a regular desktop computer instead of the powerful computer typically used by researchers. The bulk and shear modulus were computed with Reuss, Voight and Hill equations, as explained in section 2.21. The Reuss method gives the upper bound and Voight represents the lower bound, while the Hill method represents the average of both. Only the results from the Hill method are presented in this section; Reuss and Voight results appear in the appendix. Young modulus and Poisson ratios were calculated from the Hill bound.

4.6 Micro properties of the unreacted phase

Unreacted particles in the cement paste contain porosity due to grain packing. The elastic properties of C-S-H depend significantly on the scale level. For nano-scale, the structure includes nano pores, which normally have a low influence on mechanical properties. However, at micro-scale, the effect of micro pores is estimated to be relevant. The micro properties of the unreacted phase can be
calculated by using a packing factor density, which is defined by the ratio of volume of solid to total volume. Kepler (Askeland et al., 2011) proposed a packing ratio of spheres that presents the densest possible packing:

\[ \eta_{Kepler} = \frac{\pi}{\sqrt{18}} = 74.048\% \]  

(3.1)

The porosity is then calculated by:

\[ p = 1 - \eta_{Kepler} = 26\% \]  

(3.2)

One of the simplest equations used to calculate the elastic modulus (taking porosity into account) is provided by Knudsen and Helmuth (Yoshimura et al., 2007).

\[ E_{knudsen} = E_0 e^{(-kp)} \]  

(3.3)

\[ E_{Helmuth} = E_0 (1-p)^k \quad k = 3 \]  

(3.4)

Where \( E_0 \) is elastic modulus with zero porosity, \( p \) is the porosity and \( k \) fitting factor from experimental results. \( k =3.4 \) (Velez, et al., 2001)

\[ E = E_0 e^{(-3.4p)} \]  

(3.5)

Table 6 shows the Young modulus results of the clinker phases, calculated after applying the different porosities on C\(_3\)S, C\(_2\)S, C\(_3\)A and C\(_4\)AF structures.

<table>
<thead>
<tr>
<th>Clinker phase</th>
<th>C(_3)S</th>
<th>C(_2)S</th>
<th>C(_3)A</th>
<th>C(_4)AF</th>
</tr>
</thead>
<tbody>
<tr>
<td>( E_0 ) (GPa)</td>
<td>136</td>
<td>120</td>
<td>171</td>
<td>204.5</td>
</tr>
<tr>
<td>( E_{knudsen} ) (GPa)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porosity: ( p = 0.1 )</td>
<td>96.8</td>
<td>85.41</td>
<td>121.71</td>
<td>145.56</td>
</tr>
<tr>
<td>( p = 0.15 )</td>
<td>81.66</td>
<td>72.06</td>
<td>102.68</td>
<td>122.80</td>
</tr>
<tr>
<td>( p = 0.20 )</td>
<td>68.90</td>
<td>60.79</td>
<td>86.63</td>
<td>103.60</td>
</tr>
<tr>
<td>( p = 0.25 )</td>
<td>58.12</td>
<td>51.29</td>
<td>73.09</td>
<td>87.41</td>
</tr>
<tr>
<td>( p = 0.30 )</td>
<td>49.04</td>
<td>43.27</td>
<td>61.66</td>
<td>73.74</td>
</tr>
<tr>
<td>( p = 0.35 )</td>
<td>41.37</td>
<td>36.51</td>
<td>52.02</td>
<td>62.21</td>
</tr>
<tr>
<td>( p = 0.40 )</td>
<td>34.90</td>
<td>30.80</td>
<td>43.89</td>
<td>52.48</td>
</tr>
<tr>
<td>( p = 0.45 )</td>
<td>29.45</td>
<td>25.98</td>
<td>37.03</td>
<td>44.28</td>
</tr>
<tr>
<td>( p = 0.50 )</td>
<td>24.84</td>
<td>21.92</td>
<td>31.24</td>
<td>37.26</td>
</tr>
<tr>
<td>( p = 0.55 )</td>
<td>20.96</td>
<td>18.49</td>
<td>26.36</td>
<td>31.52</td>
</tr>
</tbody>
</table>
After applying porosity to the elastic modulus resulting from the molecular dynamic simulation, remarkable decrements were noticed for all clinker phases. For instance, decrements of 29%, 49%, 64%, 74% and 82% were calculated when porosity of 10%, 20%, 30%, 40% and 50% was accounted for in the different clinker phases. Figure 49 displays the impact of porosity on the Young modulus results of C₃S, C₂S, C₃A and C₄AF.

![Elastic moduli as a function of the relative porosity of the clinker phases](image)

**Figure 49:** Elastic moduli as a function of the relative porosity of clinker phases

As shown in Figure 49, there was an inverse relationship between Young modulus and relative porosity, which is represented by an exponential function. As the porosity increased, the Young modulus decreased.

### 4.7 Discussion of results

The three major phases affecting the elastic modulus of cement paste are unreacted clinker particles, C-S-H and Portlandite (CH). Unreacted cement grains can be present in the cement paste even after a long period of hydration, which can be due to the limited space for C-S-H to grow or to the low water to cement (W/C) ratio. All Young modulus results completed with molecular modeling using
COMPASS and DREIDING force fields (C2S, C3S and C3A) agreed with the experimental results reported by Velez et al. (2001) and Bass (2015). For instance, molecular modeling of C3S, C2S and C3A showed values of 136.1, 120 and 171.7, respectively, which is close to the experimental results that mention Young modulus of 135, 130, and 160, respectively. The only discrepancy that can be noted was for C4AF, in which the calculated Young modulus (204.5 GP) was almost 1.6 times higher than the experimental value (125GPa) presented in the literature. However, this disagreement has already been mentioned by several researchers (Jorge et al., 2011), (Manzano et al., 2008). For example, Jorge et al. (2011), who used an ab-initio simulation, found a noticeable difference in the elastic modulus of the ferrite phase (C4AF) (238.2 GPa) compared to the experimental results (125GPa) presented by Velez et al., (2001). According to these authors, the reasons for the disagreement between the simulation results and the experimental values is not clear. However, in regard to the different force fields used in the present study, it can be assumed that the difference may be caused by the force field used (CVFF) for C4AF. Furthermore, the disagreement in the Young modulus of the ferrite phase could also have been caused by the number of cells used in the simulation. Indeed, in all the simulations completed in this work, only one cell was used to predict elastic properties. Using a higher number of cells in molecular modeling may require a more powerful computer with longer simulation times, which is incompatible with the reduced computation simulation time targeted in this study. Although the C3S, C2S and C3A results can be considered acceptable compared to the experimental values, further investigation is suggested for C4AF.

When taking the effect of cement porosity on the elastic modulus of different clinker phases into consideration, the elastic modulus decreased as porosity increased, which is in agreement with the polynomial relationship mentioned by Powers (1960). For example, when increasing porosity from zero to 10% and 25%, the Young modulus of different clinker phases decreased from 136.1 GPa to 96.8 GPa and 68.9 GPa, respectively, which corresponds with decrements of 28.9% and 49.3%, respectively. The decrease in the estimated Young modulus is also in agreement with the experimental results presented by Velez et al. (2001), which showed decrements of about 30% to 40% when
porosity increased from 10% to 20% for C₃S, C₂S and C₃A clinker phases. The porosity results also indicate that as the packing factor of grains decreases (and porosity increases), the elastic modulus decreases considerably. This relationship between packing factor and elastic modulus is an agreement with conclusions made by Constantinides and Ulm (2007), in which they showed that material became unstable when its packing factor reached a ratio of 0.5% (relative porosity of 0.5). They also found that the elastic modulus was mainly controlled by particle-to-particle contact, which can also explain our results.

One other point from the molecular simulation is the anisotropic elastic modulus shown in all clinker phases. For example, Alite had $E_X = 145.4131$, $E_Y = 148.41$ and $E_Z = 122.61$ GPa, as shown in appendix I. This confirms that the C₃S was more compressible in Z direction ($E_Z = 122.61$) than in X and Y directions ($E_X = 145.41$ and $E_Y = 148.41$). In the experimental results presented by Velez et al. (2001) Young modulus direction was not taken into consideration, and Poisson's ratio of 0.3 was used for all constituents. However, this study shows that the low elastic modulus in Z direction can lead to higher expansion/shrinkage, shown by a high Poisson ratio $\nu_{zx}$ of 0.3869 compared to $\nu_{yx}=0.2755$ and $\nu_{yx}=0.2700$ (as shown in appendix I). This may illustrate that, in the case of C₃S, C₃A and C₂S, the results of molecular simulation can be considered more accurate than experimental results.

Another notable result was the effect of C/S ratio on the elastic modulus of clinker phases. C₃S and C₂S elastic modulus showed that the higher the C/S ratio, the higher the elastic modulus. For example, C₃S had a C/S=3 and Young modulus of 136.1 GPa, while C₂S had a C/S=2 and Young modulus of 120 GPa. These results are investigated further in the following chapters.

4.8 Summary

The aim of the work outlined in this chapter was to simulate the elastic properties of clinker phases using molecular dynamic simulation by COMPASS force field. However, this targeted force field was applicable only to C₃S and C₂S structures, which led to the use of others force fields from
Materials Studio software such as DREIDING and CVFF. The elastic properties determined by COMPASS and DREIDING force fields on C₃S, C₂S and C₃A showed agreement between the experimental values cited in literature and the estimated modulus resulting from this study. Although the CVFF molecular simulation of the C₄AF structure showed an overestimated elastic modulus compared to the experimental values, the Young modulus of C₄AF was estimated to be equal to 204.5 GPa, which is close to the results of the ab-initio simulation.

When different porosity ratios were considered, results confirmed that elastic modulus decreased as porosity increased, which is in line with the experimental results mentioned in literature.
5.1 Introduction

Portlandite Ca(OH)$_2$ is the second major hydration product of cement paste and constitutes 20-25% of its total volume. Knowledge of the elastic properties of Portlandite is important in the creation of cement paste. The presence of Portlandite in cement influences strength, reduces shrinkage and increases the alkalinity of the pore solution, which creates a passive layer that protects reinforcements from corrosion.

Ettringite is the third most relevant hydration product of Portland cement. It is a crystalline product that can affect the strength and long-term durability characteristics of concretes. The structure of Ettringite is composed of Ca$_6$[Al$_2$(OH)$_{12}$.24H$_2$O]$^{6+}$ placed on the top of each other in a C axis. One of the characteristics of Ettringite is its open structure, which allows interchange with other atoms. 80% of the atoms in Ettringite are water molecules and hydroxyl, which reduce its density to 1.7 and lead to a low elastic modulus (Speziale et al. 2008).

5.2 State of art

Elastic properties of Portlandite can be determined experimentally by X-ray spectroscopy and Brillouin zone spectroscopy. Speziale, et al. (2008) performed an experimental study on Portlandite using a natural specimen found in the Kalahari Manganese Field (South Africa). When they used Brillouin scattering measurements, they found that the Reuss and Voight bound of Portlandite were
Elastic Properties of hydration products

31.65 GPa and 20.95 GPa, respectively, and the Young modulus and Poisson ratios were 51.5 GPa and 0.23.

Aierken et al. (2015) performed a study using a Density Function Theory (DFT) and experimental tools to determine the mechanical properties and other characteristics of Ca(OH)$_2$. Due to the fact that Ca (OH)$_2$ is a layered structure, they performed these calculations on the monolayer and bilayer of calcium hydroxide. As part of the experimental work, they produced the calcium hydroxide using Ca$_3$SiO$_5$ mixed with different water-to-solid ratios in the range of 0.2 to 0.9 and heated to 40°C. The time of growth of calcium hydroxide was estimated to be 8 hours, with a water-to-solid ratio of 0.6 and a synthesized crystal dimension of 0.1-2 mm. To calculate the mechanical properties, they stacked these layers on the top of each other. The Young modulus of the calcium hydroxide was estimated to be 50.7 GPa and 55.0 GPa for the monolayer and bulk, respectively, whereas the shear modulus was 19.16 GPa and 21.23 GPa for the monolayer and bulk, respectively.

Speziale et al. (2008) also performed an experimental study using Brillouin spectroscopy to determine the elastic properties of Ettringite. They found the Young modulus was 25 GPa, the Hill average for the bulk modulus was 27.15 GPa and the Hill average of shear modulus was 9.5 GPa. They concluded that the elastic tensor is strongly anisotropic, which relates to the lattice structure of mineral computation.

5.3 Computational Method

5.3.1 Lattice parameters

The structures of Portlandite and Ettringite used in this study were previously resolved by Henderson and Gutowsky (1962) and by Hartman and Berliner (2006), respectively. The lattice parameters ($a$, $b$, $c$, $\alpha$, $\beta$, and $\gamma$) reported by those authors and used in the simulation of this study are presented in Table 7.
Table 7: Lattice parameters for Portlandite and Ettringite

<table>
<thead>
<tr>
<th>Lattice Parameters</th>
<th>CH</th>
<th>Ettringite</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>11.6389</td>
<td>11.260</td>
</tr>
<tr>
<td>b</td>
<td>14.1716</td>
<td>11.260</td>
</tr>
<tr>
<td>c</td>
<td>13.6434</td>
<td>21.480</td>
</tr>
<tr>
<td>α</td>
<td>104.982</td>
<td>90.000</td>
</tr>
<tr>
<td>β</td>
<td>94.622</td>
<td>90.000</td>
</tr>
<tr>
<td>γ</td>
<td>90.107</td>
<td>120.000</td>
</tr>
<tr>
<td>Ref</td>
<td>[a]</td>
<td>[b]</td>
</tr>
</tbody>
</table>

[a]: (Henderson and Gutowsky, 1962)  
[b]: (Hartman and Berliner, 2006)

The first step in our simulation was to build the crystal structure. To do so, the parameters mentioned in Table 7 were incorporated into the model data. Then, the atoms were added to the crystal depending on their types and coordinates.

5.3.2 Geometry optimization parameters

After trying COMPASS force field on both structures of Portlandite and Ettringite, it was decided that this type of molecular dynamic simulation is not applicable to the structure of Ettringite. For this reason, two force fields were used in this simulation: COMPASS force field for Portlandite and Universal for Ettringite. Table 8 shows the final parameters used to optimize the lattice structure.

Table 8: Geometry optimization parameters

<table>
<thead>
<tr>
<th>Geometry minimization parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Force field</td>
</tr>
<tr>
<td>Quality</td>
</tr>
<tr>
<td>Summation Method</td>
</tr>
<tr>
<td>Repulsive cutoff</td>
</tr>
</tbody>
</table>

5.3.3 Dynamic optimization parameters

Table 9 shows the parameters used in the dynamic simulation of Portlandite and Ettringite:
Table 9: Dynamic optimization parameters

<table>
<thead>
<tr>
<th>Dynamic Simulation parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ensemble</td>
</tr>
<tr>
<td>Temperature</td>
</tr>
<tr>
<td>Pressure (GPa)</td>
</tr>
<tr>
<td>Time step (fs)</td>
</tr>
<tr>
<td>Total simulation (ps)</td>
</tr>
<tr>
<td>Thermostat</td>
</tr>
<tr>
<td>Barostat</td>
</tr>
</tbody>
</table>

After minimizing the structure, the data mentioned in Table 9 were incorporated. The dynamic simulation was run in which the atoms interacted with each other for a period of time. The position of atoms was determined by numerically solving the Newton equation, whereas forces between atoms were calculated with the potential energy using different force fields.

5.4 Dynamic simulation results

When the different forces between atoms are calculated, the mechanical properties of the crystals can be determined. Table 10 shows the mechanical properties of Portlandite and Ettringite resulting from the simulation in this study. For comparison purposes, some experimental results mentioned in literature are also presented in Table 10.

Table 10: MD simulation results of Portlandite and Ettringite (Appendix VI)

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Crystals</td>
<td>CH</td>
<td>Ettringite</td>
<td></td>
</tr>
<tr>
<td>Bulk Modulus :K(GPa)</td>
<td>46.3</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Shear modulus: G(GPa)</td>
<td>17.8</td>
<td>7.33</td>
<td></td>
</tr>
<tr>
<td>Young modulus: E(GPa)</td>
<td><strong>47.3</strong></td>
<td><strong>18.9</strong></td>
<td></td>
</tr>
<tr>
<td>ν : Poisson ratio</td>
<td>0.329</td>
<td>0.29</td>
<td></td>
</tr>
<tr>
<td>Force field used</td>
<td>FC</td>
<td>FU</td>
<td></td>
</tr>
<tr>
<td>Ref E (GPa)</td>
<td>51.35[^c]</td>
<td>55[^d]</td>
<td>25[^e]</td>
</tr>
</tbody>
</table>
The bulk modulus, shear modulus, Young modulus and Poisson ratio were obtained after running the dynamic simulation, as shown in Table 10. Those mechanical characteristics were the final results targeted in our simulation.

5.5 Discussions

After minimizing the structure of calcium hydroxide and Ettringite, the calculation of elastic modulus reported in this section was carried out by molecular dynamic simulation, using COMPASS and Universal force fields in Materials Studio software. Young and shear modulus estimated for Portlandite (47.3 GPa and 17.8 GPa, respectively) were slightly lower than the experimental results reported by Speziale et al. (2008) (which were 51.35 GPa and 20.95GPa), who used Brillouin spectroscopy (section 4.1). In addition, the average bulk modulus found in this study (46.3GPa) was slightly higher than the result found by Speziale et al. (2008), which was 31.65 GPa. This small inconsistency can be related to the accuracy of the experimental tools used to measure the elastic properties, in which the porosity may not have been accounted for. Another reason may be that COMPASS force field was used to determine Portlandite structure. It is well known from the literature that the type of force field is the principal parameter affecting the results of the molecular dynamic simulation.

Although small differences were noted compared to the experimental results mentioned in literature, the results of Portlandite agreed with the elastic properties found by Aierken et al. (2015), who used the first principle calculation method. For example, Aierken et al. (2015) calculated Young modulus of 55GPa, bulk modulus of 44 GPa, shear modulus of 21.23 GPa and Poisson ratio of 0.30, which are close to the results of this study.
Ettringite results shown in Table 11 indicate bulk, shear and Young modulus of 15 GPa, 7.33 GPa, 10 and 18.9 GPa, respectively. These values are slightly lower than the experimental results completed by Speziale et al. (2008), which showed bulk, shear and Young modulus of 27.15 GPa, 9.5 GPa and 25 GPa, respectively. However, the results of the current study agree with the other simulation methods shown in literature; for instance, Kamali et al. (2004) determined elastic modulus of 22.4 GPa and a Poisson ratio of 0.25 by using the ELAS3D modeling program.

When considering the structure proposed by Hartman and Berliner (2006) used in this study, Ettringite is composed of \([\text{Al(OH)}_6]^{3-}\) and water molecules with sulfate ions. As shown in Figure 31, section 1.13.2 (Chapter 1), \([\text{Al(OH)}_6]^{3-}\) linked through \(\text{Ca}^{2+}\) run parallel to c axis, while water molecules with sulfate ions are in the inner part of the structure. In the Ettringite formula \([\text{Ca}_3\text{Al(OH)}_612\text{H}_2\text{O}_2 (\text{SO}_4)3.2\text{H}_2\text{O}]\), most of the atoms are water molecules or hydroxyl, which result in a reduced Ettringite density estimated at 1.79 (Materials Studiov7). This result agrees with the experimental density of Ettringite (1.7) mentioned in section 4.2. The lower density of Ettringite can justify its low elastic modulus, estimated by using molecular dynamic simulation.

The simulation results (shown in Appendix VI) indicate that the Young modulus of Ettringite in the three axes are: \(E_x=14.03\) GPa, \(E_y=15.8\) GPa and \(E_z=32.6\) GPa, which is anisotropic. The structure in z-axis represents the stiffer direction compared with x and y directions. The Young modulus in z-axis (\(E_z\)) represents values of around two times higher than the other axes (\(E_x\) and \(E_y\)), which is in good agreement with results of Speziale et al. (2008), which showed \(E_z\) to be equal to 1.6 times \(E_x\). The crystallography form of Ettringite composed of \([\text{Al(OH)}_6]^{3-}\) can be the reason of its stiffer structure in c-axis (z direction).

A comparison between the Portlandite and Ettringite results shows that the Young modulus of Ettringite is around 60% lower than that of the Portlandite, which indicates that Ettringite stiffness is around 2.5 times lower than that of Portlandite. The low stiffness of Ettringite is due to a structure composed mostly of water molecules, which decrease its density to 1.79 g.cm\(^3\). It is well known that
density is one of the paramount factors affecting the mechanical properties of materials. The water in Ettringite structure reduces the cohesion between particles and shields the interatomic forces.

The Portlandite investigated in this study supports the use of COMPASS force field. However, this method of simulation in not applicable for Ettringite structure. COMPASS force field is not designed for cementitious materials, so the low Young modulus of Ettringite can be attributed to the precision and accuracy of COMPASS force field used for Portlandite compared to the Universal force field used for Ettringite.

5.6 Summary

The objective of the work detailed in this chapter has been to determine the elastic modulus of Portlandite and Ettringite. The molecular dynamic simulation was carried out with two force fields: COMPASS and Universal. Compared to the experimental results shown in literature, the elastic properties of Portlandite are slightly different. However, good agreement was noticed between the simulation results of this study and others computational methods.

Like Portlandite, the results of Ettringite also show differences compared to the experimental results and are in close agreement with the results of other simulation methods (e.g. Density Function Theory (DFT)). In addition, the Young modulus of Ettringite shows anisotropy in which the z-direction value is twice that of the two other directions. The c-axis (z) represents the stiffer axis due to the structure of Ettringite that runs in z- axis with $[\text{Al(OH)}_6]^{3-}$.

In comparing results of Portlandite and Ettringite, a higher elastic modulus of Portlandite was noticed, which is related to the high water molecules in the structure of Ettringite compared to Portlandite.
Chapter 6  Elastic Properties of analog minerals

6.1 Introduction

Although there are 30 mineral crystals with chemical composition similar to C-S-H created from cement paste, there are some differences in atomic arrangement, Ca/Si ratio and the number of OH and H₂O (Taylor, 1997). According to literature, Tobermorite and Jennite are the minerals with the most similarities to C-S-H. There are three types of Tobermorite: Tobermorite 9Å, Tobermorite 11Å and Tobermorite 14Å, which differ by their degree of hydration. Jennite is also similar to C-S-H structure at later ages of hydration, with a Ca/Si ratio of 1.5 (as explained in chapter 1, section 1.11.4). The purpose of this chapter is to determine the elastic modulus of analog C-S-H minerals such as the Tobermorite family and Jennite with the use of molecular dynamic simulation by COMPASS force field. It also compares Young modulus results from this study with previous experimental and theoretical results in literature. This final purpose of this chapter is to determine the Tobermorite and Jennite structure closest to C-S-H.

6.2 State of art of elastic properties of Tobermorite and Jennite

6.2.1 Tobermorite 14Å

Oh et al. (2012) performed an experimental study to determine the bulk modulus of Tobermorite 14Å. They measured the unit cell volume change by increasing the pressure to 4.8 GPa using high-pressure synchrotron X-ray diffraction under hydrostatic condition. The Tobermorite sample was finely ground and mixed with a pressure transmitting liquid (methanol/ethanol) to generate hydrostatic pressure. The X-ray diffraction pattern was collected at different pressures. As the applied hydrostatic
pressure was increased, the unit cell of Tobermorite 14 Å deformed, leading to a change in the lattice parameters. By analyzing X-ray diffraction patterns at different pressures, the lattice parameter was determined based on the positions of diffraction peaks. The volume of the lattice was calculated from the lattice parameters. Only the bulk modulus of Tobermorite 14A was determined to be $K_0 = 47$ GPa. They also found that the bulk modulus of Tobermorite 14A depends on lattice parameter c, which is related to the interlayer spacing composed of dreierketten silicate chains, interlayer Ca, and water molecules (Oh et al., 2012).

Among the different theoretical approaches, molecular dynamics and ab-initio methods were the most common techniques used in literature to determine the elastic properties of Tobermorite 14A. The theoretical results show some differences, which depend on simulation method, number of cells and force field used.

6.2.2 Tobermorite 11Å

The Tobermorite 14Å was heated at a temperature of 125°C for 24 hours, which led to the formation of Tobermorite 11Å with a peak diffraction around 11.3Å, which also indicates reduced interlayer thickness. Using the same experimental method described in section 5.2.1 (chapter 5), Moon (2013) calculated the bulk modulus of Tobermorite 11Å to be $K_0 = 62 (4)$ GPa where the value between parentheses represents the standard deviation.

6.2.3 Tobermorite 9Å

Moon (2013) performed an experimental study on Tobermorite 9Å to determine its bulk modulus. The Tobermorite 9Å was prepared by heating Tobermorite 14Å at 400°C. The bulk modulus of Tobermorite 9Å was determined by measuring the unit cell volume change by increasing the pressure up to 8.5 GPa using high-pressure synchrotron X-ray diffraction under a hydrostatic condition, using the same procedure described for Tobermorite 14Å. Moon (2013) calculated the bulk modulus of Tobermorite 9Å to be $K_0 = 82 (4)$ GPa.
6.2.4 Jennite

Jennite is the other mineral that is believed to be analog to C-S-H at a late stage of hydration. The structure of Jennite was solved by Bonnacorsi et al. (2004). Moon et al. (2015) performed an experimental study on Jennite to determine its bulk modulus, and also used the first principal method to determine its other elastic properties. Using high-pressure synchrotron X-ray diffraction under hydrostatic, Moon et al. (2015) found that the bulk modulus of Jennite is \( K_0 = 64 \pm 2 \) GPa, where the value between parentheses represents the standard deviation. From the first principal method, they found bulk, shear and Young modulus were 62 GPa, 28 GPa and 72.9 GPa, respectively, with a Poisson ratio of 0.3.

6.3 Computational method

6.3.1 Crystal parameters

As the first step in the molecular simulation, the crystal structure was built by entering the data of the space group (which represents the symmetry of the structure), lattice type (triclinic or monoclinic) and lattice parameters \((a, b, c, \alpha, \beta, \text{and } \gamma)\) of Tobermorite 9A, 11A, 14A and Jennite. Those data (summarized in Table 11) were resolved experimentally by Merlino et al. (1999 and 2001) and Bonaccorsi et al. (2004 and 2005).
<table>
<thead>
<tr>
<th>Lattice Parameters</th>
<th>Tobermorite 9A</th>
<th>Tobermorite 11A</th>
<th>Tobermorite 14A</th>
<th>Jennite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Space group</td>
<td>C1</td>
<td>BM</td>
<td>B11b</td>
<td>P1</td>
</tr>
<tr>
<td>Lattice type</td>
<td>Triclinic</td>
<td>Monoclinic</td>
<td>Monoclinic</td>
<td>Triclinic</td>
</tr>
<tr>
<td>a</td>
<td>11.156</td>
<td>6.735</td>
<td>6.735</td>
<td>10.576</td>
</tr>
<tr>
<td>b</td>
<td>7.303</td>
<td>7.385</td>
<td>7.425</td>
<td>7.265</td>
</tr>
<tr>
<td>c</td>
<td>9.566</td>
<td>22.487</td>
<td>27.987</td>
<td>10.931</td>
</tr>
<tr>
<td>α</td>
<td>101.080</td>
<td>90.000</td>
<td>90.000</td>
<td>101.300</td>
</tr>
<tr>
<td>β</td>
<td>92.830</td>
<td>90.000</td>
<td>90.000</td>
<td>96.980</td>
</tr>
<tr>
<td>γ</td>
<td>89.980</td>
<td>123.250</td>
<td>123.250</td>
<td>109.650</td>
</tr>
<tr>
<td>Ref</td>
<td>[a]</td>
<td>[b]</td>
<td>[c]</td>
<td>[d]</td>
</tr>
</tbody>
</table>

[a]: (Merlino et al., 1999)  
[b]: (Merlino et al., 2001)  
[c]: (Bonaccorsi et al., 2005)  
[d]: (Bonaccorsi et al., 2004)

As shown in Table 11, Tobermorite 9A, 11A, 14A and Jennite represent different space groups. Although Tobermorite 9A and Jennite, and Tobermorite 11A and 14A have the same triclinic or monoclinic lattice types, respectively, these similarities in crystal family (lattice type) have not been reported in literature, offering no preliminary indication or conclusion. After building the crystal structure using the data in Table 11, the different types of atoms were added to the crystal unit cell, which represents an imaginary parallelepiped of a repeating pattern.

6.3.1.1 Geometry optimization parameters

When the final structures of Tobermorite 9A, 11A, 14A and Jennite are built with atoms, a geometry optimization is usually needed to bring the geometry to a stable state. The process of geometry optimization or minimization is an iterative process where the energy structure is brought to its minimum energy and the force between atoms is zero. More information about the geometry optimization in the molecular simulation is provided in chapter 2, section 2.12. Table 12 shows the
different parameters used to optimize the lattice structure including force field, quality, summation method and cut-off distance.

Table 12: Geometry minimization parameters

<table>
<thead>
<tr>
<th>Geometry minimization parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Force field</td>
</tr>
<tr>
<td>Quality</td>
</tr>
<tr>
<td>Summation Method</td>
</tr>
<tr>
<td>Repulsive cutoff</td>
</tr>
</tbody>
</table>

The appropriate force field is very important in molecular dynamic simulation, since it determines the potential energy function of the structure, and adjusts atom position to reduce energy expression by using different algorithms to locate the minimum energy. In this study, COMPASS force field was applicable for all the analog C-S-H minerals.

The calculation of minimum energy was controlled by the quality setting. Ultrafine quality was used because it offers the most precise energy calculation. For instance, energy accuracy is set to $2 \times 10^{-5}$ kcal/mol for ultrafine quality, while it is set to 0.02 kcal/mol for coarse quality. The Ewald-type summation method was used in this study because it has been reported in literature as the best option for the calculation of non-bond energy in periodic lattice. A cut-off distance of 12.5Å was used to reduce the number of interactions and to cut the contribution of van der Waals energy for Ewald summation.

6.3.1.2 Dynamic simulations parameters

The dynamic simulation was performed after optimizing the structure. The crystal system was simulated as it occurs in the natural environment, where it may be under external pressure and varied temperatures. Table 13 shows the different parameters used in the molecular dynamic simulations of the present study.
Table 13: MD simulation parameters

<table>
<thead>
<tr>
<th>Dynamic Simulation parameters</th>
<th>NPT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ensemble</td>
<td>NPT</td>
</tr>
<tr>
<td>Temperature</td>
<td>298 K (25°C)</td>
</tr>
<tr>
<td>Pressure (GPa)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Time step (fs)</td>
<td>0.5</td>
</tr>
<tr>
<td>Total simulation (ps)</td>
<td>200-500</td>
</tr>
<tr>
<td>Thermostat</td>
<td>Nose</td>
</tr>
<tr>
<td>Barostat</td>
<td>Anderson</td>
</tr>
</tbody>
</table>

Different methods can be used to control temperature and pressure. For all the simulations in this study, an NPT ensemble was used (where N is number of particles, P is pressure and T is temperature), which remained fixed where the unit cell was allowed to change. (Other ensembles such as NVE and NVT can be also used.) Thermostat and Barostat are temperature and pressure control methods used to regulate temperature and pressure, respectively. One of the most important parameters is the time step integration of Newton equation, where higher time steps can lead to errors in the simulation. Additional information about ensembles is provided in chapter 2, section 2.19.

6.4 Results of molecular simulation

6.4.1 Elastic properties of Tobermorite and Jennite

Tables 15 and 16 show the elastic modulus of Tobermorite 9Å, 11Å, 14Å and Jennite resulting from our simulation using COMPASS force field. Other experimental and theoretical results from literature are also presented in Tables 14 and 15 for comparison.
Table 14: Elastic properties of Tobermorite 9Å and Tobermorite 11Å (Appendix VII, VIII)

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Tobermorite 9Å</th>
<th>Tobermorite 11Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_0$ (GPa)</td>
<td>87.7</td>
<td>48.3</td>
</tr>
<tr>
<td>$G$ (GPa)</td>
<td>34.7</td>
<td>25.7</td>
</tr>
<tr>
<td>$E$ (GPa)</td>
<td>92</td>
<td>65.5</td>
</tr>
<tr>
<td>$\nu$</td>
<td>0.34</td>
<td>0.27</td>
</tr>
</tbody>
</table>

Ref : $K_0$ (GPa): 71.42[e], 68.95[f], 86[g](70-82)[g] 66.65[e], 73.57[f], 68[g], 67[g]
$G$ (GPa): 37.11[e], 37.44[f], (50.3-54.1)[g] 32.03[e], 29.19[f], 32[g]
$E$ (GPa): 95.06[e], 95.11[f], (122.9-130.3)[g] 82.82[e], 77.34[f], 83.1[e]

Table 15: Elastic properties of Tobermorite 14Å and Jennite (Appendix IX, X)

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Tobermorite 14Å</th>
<th>Jennite</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_0$ (GPa)</td>
<td>40.9</td>
<td>73.3</td>
</tr>
<tr>
<td>$G$ (GPa)</td>
<td>16.7</td>
<td>23.9</td>
</tr>
<tr>
<td>$E$ (GPa)</td>
<td>44.1</td>
<td>64.7</td>
</tr>
<tr>
<td>$\nu$</td>
<td>0.32</td>
<td>0.35</td>
</tr>
</tbody>
</table>

Ref : $K_0$ (GPa): 35.91[e], 44.8[f], 44[g], 47, 47[h] 31.83[e], 40.2[f], 64[i], 62[i]
$G$ (GPa): 20.61[e], 19[f], 41[g] 21.96[e], 22.10[f], 28[i]
$E$ (GPa): 51.9[e], 49.94[f], 99.5[g] 53.55[e], 56.03[f], 72.9[i]

Ko: Bulk modulus
G: shear modulus
E: Young modulus
$\nu$: Poisson ratio

[e]: (Shahsavari et al., 2009)
[f]: (Manzano et al., 2007)
[g]: (Moon, 2013) (experimental and computation)
[h]: (Oh et al., 2012)
[i]: (Moon 2015) (experimental and computation)

As seen from Table 14, the Young modulus of Tobermorite 9Å in this study was 92GPa, which is in good agreement with the results of Shahsavari et al. (2009) and Manzano et al. (2007); they showed a modulus of around 95 GPa using other molecular dynamic simulation method. However, the experimental method performed by Moon (2013) showed slightly higher results that ranged between 122.9 GPa and 130.3 GPa.

Although the Young modulus of Tobermorite 11Å was estimated to be equal 65.5GPa, the simulation in this study showed it to be slightly lower than the theoretical and experimental results attained by
Shahsavari et al. (2009) (82.82 GPa) and Moon (2013) (83.1 GPa). Manzano et al. (2007) found a modulus of 77.34 GPa, which is only 11% higher than our results. In addition, the difference between the experimental values found by Moon (2013) and the Young modulus of our simulation do not exceed 21%, which is acceptable compared to other simulation methods.

The Young modulus of Tobermorite 14A, resulting from our molecular dynamic simulation with COMPASS force field, was estimated to be 44.1 GPa, which is close to the modulus found by Manzano et al. (2007) (49.94 GPa) using different molecular dynamic method. Although the experimental value showed by Moon (2007) was significantly higher than our simulation result, it was also higher than the other numerical simulation results completed by Shahsavari et al. (2009) and Manzano et al., (2007). This indicates that all numerical simulation methods underestimate the Young modulus of Tobermorite 14 A.

The elastic properties of Jennite displayed in Table 16 show the Young modulus estimated in this study to be 64.7 GPa. Although this value is slightly higher than those expected by the ab-initio numerical method completed by Shahsavari et al. (2009) (53.55 GPa) and Manzano et al. (2007) (56.03 GPa), it is closer to the result measured by Moon (2013). The difference between the Young modulus found by COMPASS force field (in this study) and the experimental value (Moon result) is around 11%, which is two times lower than the difference between the other theoretical results completed by the ab-initio method and the experimental Young modulus found by Moon (2013).

6.4.2 Relation between Young modulus and interlayer spacing

Figure 50 present the variation of elastic modulus results as a function of the interlayer spacing
Figure 50 shows that the interlayer spacing considerably affects the elastic modulus of the Tobermorite family (9A, 11A and 14A). For instance, Tobermorite 9A (characterized by its thinner interlayer spacing) has a higher elastic modulus, whereas 14A (with its larger interlayer spacing) is the softest Tobermorite mineral. It is clear from Figure 50 that the increase in interlayer spacing causes the elastic modulus to decrease.

6.4.3 Relation between Young modulus and C/S, density and water

Table 16 presents some important parameters of Tobermorite 9A, 11A, 14A and Jennite, such as C/S ratio, water to calcium ratio (W/Ca), water to silica ratio (W/Si) and density. C/S, W/Ca, W/Si were determined from different chemical formulas, whereas density values were extracted from the materials studio software before bringing the structure to its relaxed state, and before running the molecular dynamic (MD) simulation. The relationships between elastic modulus and C/S, density, W/Ca and W/Si ratios are presented in Figures 51, 52, 53 and 54, respectively.

Table 16: Properties of different types of Tobermorite and Jennite (C/S, density, W/Ca and W/Si and elastic properties)
### Elastic Properties of analog minerals

<table>
<thead>
<tr>
<th>Mineral</th>
<th>C/S</th>
<th>Density</th>
<th>W/Ca</th>
<th>W/Si</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tobermorite 9Å</td>
<td>0.83(5/6)</td>
<td>2.681</td>
<td>0</td>
<td>0</td>
<td>95.23</td>
</tr>
<tr>
<td>Ca$_5$Si$<em>6$O$</em>{16}$(OH)$_2$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tobermorite 11 Å</td>
<td>0.67(4/6)</td>
<td>2.453</td>
<td>1.25</td>
<td>0.83</td>
<td>65.5</td>
</tr>
<tr>
<td>Ca$_4$[Si$<em>6$O$</em>{15}$(OH)$_2$] · 5H$_2$O</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tobermorite 14 Å</td>
<td>0.83(5/6)</td>
<td>2.245</td>
<td>1.4</td>
<td>1.16</td>
<td>44.1</td>
</tr>
<tr>
<td>Ca$_5$[Si$<em>6$O$</em>{16}$(OH)$_2$]. 7H$_2$O</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jennite</td>
<td>1.5(9/6)</td>
<td>2.32</td>
<td>0.89</td>
<td>1.33</td>
<td>64.7</td>
</tr>
<tr>
<td>Ca$_9$(Si$<em>6$O$</em>{18}$(H$_2$)(OH)$_6$ 8H$_2$O</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

As shown in Figure 51, there is no clear relationship between C/S ratio and Young modulus results of analog C-S-H minerals. For example, while Tobermorite 9Å and 14Å had the same C/S ratio of...
0.83, they also show diverse elastic modulus. In addition, Tobermorite 11Å and Jennite have close elastic modulus results, although they present large differences in C/S ratio, with values of 0.67 and 1.5, respectively. Figure 52, however, indicates a clear relationship between density and the difference in elastic modulus of the Tobermorite family and Jennite. Tobermorite 9A, which showed the highest elastic modulus, has the higher density. Tobermorite 14A, with the lowest density, is characterized by its lower elastic modulus.

Water has a direct impact on elastic modulus results. Figures 53 and 54 show that as interlayer water increased by Ca or Si, elastic modulus decreased. The Tobermorite 14A with the highest W/Ca and W/Si ratios also had the lowest modulus of elasticity. This can be explained by the fact that, as water in the interlayer spacing increased, the silicate chains became more distant from each other and reduced the interlayer interaction between atoms in the molecular dynamic simulation, as well as the cohesiveness of the structure. In addition, the reduced interlayer spacing of Tobermorite 9A and 11A creates a denser structure than in 14A, which is also related to the quantity of water in the interlayer spacing and hence the elastic modulus.

6.5 Discussion of results

Molecular dynamics simulations using COMPASS force field were performed to determine the elastic properties of analog C-S-H minerals (Tobermorite and Jennite). The values resulting from those simulations had similarities and differences to the results of methods previously used in literature. The greatest differences were evident with the ab-initio method, which can be explained by the difference in the number of cells used in each dynamic simulation method and in the force field used. This explanation is in line with conclusions made by Wu et al. (2011), who investigated the combination of different simulation cell sizes and force fields. These authors confirmed that simulation is largely dependent on force field and cell size. Another reason may be the position of hydrogen in the water molecule. Since no information was provided about hydrogen coordinates in
the other studies, its position can be very different from one simulation to another, which may also influence results.

The elastic modulus of Tobermorite 9Å was higher than in 11Å and 14Å. The dissimilarity in elastic modulus between the different types of Tobermorite can be explained by the difference in interlayer thickness, in which Tobermorite 9Å has the thinner interlayer, and 14Å has the larger interlayer. This explanation agrees with conclusions made by Oh et al. (2012), who used X-ray diffraction to determine the bulk modulus of Tobermorite 14Å. They stated that modulus values of Tobermorite 14Å are controlled by the incompressibility of the lattice parameter c (z-axis), which is parallel to the interlayer spacing. The increase in the interlayer spacing reduces the cohesion between silicates chain, which leads to the decrease in elastic modulus.

Based on the density values of Tobermorite cells (9Å, 11Å, and 14Å) and Jennite, Tobermorite 9Å structure has the highest density with the highest elastic modulus, and Tobermorite 14Å has the lowest density with the lowest elastic modulus. The clear relationship between elastic modulus and density can be related to the packing of particles. As density increases, particles get closer to each other, which also increases elastic modulus. This suggestion agrees with the results of Constantinides and Ulm (2007), which showed a correlation between low density and low stiffness.

As shown in the correlation between elastic modulus and W/Ca, and W/Si, when water increases, Young modulus decreases. This conclusion is in line with most of the results presented in literature. According to Shahsavari et al. (2009), when excess water molecules enter the interlayer, the interlayer space increases and Coulombic interactions reduce. This can cause reduction of the elastic properties in the interlayer direction as W/Ca or W/Si increases.

Another important point concerns the results of Young modulus estimated in the three axes (Appendices VII, VIII, IX and X). With Young modulus results of Ex=86.74GPa, Ey=137.26GPa, and Ez=110.4GPa, Tobermorite 9Å, which did not contain any water molecules, showed anisotropic properties due to its layered structure. The highest value was in the y-axis (b) direction, parallel to the silicate chain, which means that the silicate tetrahedral SiO₄ had more of an effect on
determination of Young modulus. This conclusion agrees with the results found by Tunega end Zaoui (2010), which showed that the stiffest Young modulus is on the y-axis (b) direction, and is related to the silicate chain that runs in that direction.

Tobermorite 11A, which contained some water molecules in the interlayer, presented elastic modulus of Ex= 65.19GPa, Ey=94.62GPa and Ez= 67.19 GPa (Appendix VIII) in the three axes. These results also confirm an anisotropy, in which the elastic modulus in x and y-direction were similar (Ex~Ez), whereas the strongest Young modulus was on the y-axis parallel to the silicate chain. These findings also confirm the Tunega end Zaoui (2010) results mentioned above.

As mentioned in Chapter 1, section 1.10, the Young modulus of C-S-H (LD and HD) measured experimentally by nano-indentation, ranges between 20 and 30GPa, as noted by Canstantinides et al. (2013). However, in this study, the Young modulus results of Tobermorite 9A, 11A, 14A and Jennite were found to be equal to 95.23 GPa, 65.5 GPa, 44.1GPa and 64.7 GPa, respectively. This shows that the results of the analog C-S-H minerals present higher values than the experimental Young modulus of C-S-H measured by nano-intendation. The greater differences between the results of our simulation and the Young modulus of C-S-H measured experimentally can be explained by two reasons: 1) the infinite silica chain length used in our molecular simulation, or 2) the fact that the porosity of different analogue C-S-H minerals was not accounted for. These suggestions are covered in more detail in Chapter 7.

6.6 Summary

The purpose of this chapter was to evaluate the use of molecular dynamic simulation by COMPASS force field to estimate the elastic modulus of different types of Tobermorite (9A, 11A and 14A) and Jennite, which are assumed to have some similarities with the structure of C-S-H. Parameters that affect the elastic modulus variations were also determined.

The results achieved in this study present some dissimilarities with the experimental measurements, especially for Tobermorite 9A and 14A. However, most of the Young modulus results line up with
the theoretical values reported in literature. In addition, the Young modulus of Tobermorite 11A and Jennite present only small differences (around 11%) compared to the experimental measurements outlined in literature.

Based on the results of this study, the parameters that considerably affect the elastic properties of analog C-S-H minerals are: 1) interlayer spacing for different types of Tobermorite, 2) W/Ca and W/Si, and 3) density. For the first parameter (interlayer spacing), as spacing was reduced, elastic modulus increased and the structure became more compressible and less stiff. For the second parameter (W/Ca and W/Si), excess water molecules negatively affected elastic modulus by increasing interlayer spacing and reducing the interaction force between layers. The third parameter (density) is related mostly to the packing of particles, in which elastic modulus increased when density increased.

Although all analog C-S-H minerals showed higher Young modulus compared to C-S-H, Tobermorite 9A presented the highest difference, with a value more than three times that of C-S-H. In addition, the comparison between C/S ratios of C-S-H (LD and HD), measured experimentally, (which ranged between 1.2 and 2.3, as discussed in chapter 1, section 1.7) and those of Tobermorite 9A (0.83), 11A (0.67), 14A (0.83) and Jennite (1.5), showed that Tobermorite 11A had the highest difference compared to C-S-H. Based on these results, it can be concluded that Tobermorite 14A and Jennite are the minerals closest to C-S-H. For that reason, only Tobermorite 14A and Jennite are considered in Chapter 7.
Chapter 7  C-S-H gel Elastic properties

7.1 Introduction

In chapter 6, the Young modulus values of Tobermorite 9Å, 11Å, 14Å and Jennite crystals were remarkably higher than those in the experimentally measured C-S-H gel. When the Young modulus values and C/S ratios of Tobermorite and Jennite are compared to those of C-S-H, Tobermorite 14Å and Jennite are the closest minerals to C-S-H in terms of structure. However, with values of 44.1GPa for Tobermorite 14Å and 64.7 GPa for Jennite, those analog C-S-H minerals have a Young modulus that is around two times higher than that of C-S-H, which ranges between 20 and 30GPa. The goal of this chapter is to investigate the use of molecular dynamic simulation with COMPASS force field to reduce the Young modulus of Tobermorite 14Å and Jennite and make it closer to that of C-S-H.

Two methods were considered:

The first method is based on conclusions made by Taylor (1997). In that study, the structure of C-S-H showed similarities with Tobermorite and Jennite, with finite silicate chain following the rule of 3n-1 (where n is an integer number). The structure of Tobermorite 14Å and Jennite in our molecular simulation were modified to be finite silicate chain in which n=1 (dimer), rather than the infinite chain used in chapter 6.

In the second method, because the experimental elastic modulus of C-S-H covered in literature was measured in low and high density (LD and HD) states with porosity equal to 36% and 26%, respectively, and since the porosity of different analog C-S-H minerals was not accounted for in our previous simulations, this important parameter was investigated for Tobermorite 14Å and Jennite with infinite and finite silica chains.
In addition, the results of this study and the experimental elastic modulus of C-S-H mentioned in literature were compared to verify the reliability of the proposed solutions.

7.2 State of art on finite silicate chain

The omission of the bridging tetrahedral silicon in Jennite does not change the total charge of the unit cell, where one silicon atom has +4e⁺ and two atoms of oxygen (-2e⁻) are omitted. In the case of Tobermorite, the removal of one silicon atom (4e⁺), one oxygen (-2e⁻) and one hydroxyl (-1e⁻) leads to a negatively charged system (Figure 55). Taylor (1997) suggests that a hydrogen atom be added to the oxygen atom to equilibrate the system.

![Figure 55: Tobermorite structure with infinite and finite chain (bridging silicate removed)](image)

Because the silicate chain length has a direct relationship with C/S ratio, removing the bridging silicate chain also modifies the C/S ratios of Tobermorite 14A and Jennite. Omitting the bridging silicate tetrahedral is equivalent to removing SiO₂ from the structure; the hydrogen in the silanol group is moved to the pair silicate tetrahedral. Consequently, the removal of SiO₂ increases the C/S ratio of Tobermorite 14A and Jennite. Indeed, according to the atomic model proposed by Taylor (1997) (chapter 2, section 2.7.4), the removal of the bridging increases C/S ratios of Tobermorite 14A and Jennite from 0.83 to 1.25 and from 1.5 to 2.25, respectively. These new C/S ratios of Tobermorite 14A and Jennite with modified chain are in the range of C/S ratios of C-S-H measured experimentally.
to be between 1.2 and 2.3, which suggests that the molecular dynamic simulation of modified structures of Tobermorite 14A and Jennite can result in an elastic modulus closer to that of C-S-H. Limited studies have been conducted into the simulation of defect structures of Tobermorite and Jennite, with only the ab-initio method used in molecular simulation. For instance, Manzano et al. (2007) completed a molecular simulation by ab-initio method on modified Tobermorite 14A and Jennite structures. They found the Young modulus to be equal to 35.6 and 43.7 GPa for Tobermorite 14A and Jennite, respectively, which is close to the Young modulus of C-S-H. No molecular dynamic simulation has been done using COMPASS force field to determine the elastic properties of Tobermorite 14A and Jennite with modified structure.

In this study, the defect Tobermorite 14A and Jennite are called T2 and J2, respectively, indicating a silicate chain without a bridging silicate.

7.3 Computational method of finite silicate chain of Tobermorite 14A and Jennite

7.3.1 Geometry optimization parameters

Geometry optimization of molecules brings the structure to its stable state. As a first step in all molecular dynamic simulations using COMPASS force field presented in this chapter, geometry optimization was run to bring the structure of modified Tobermorite 14A (T2) and Jennite (J2) to its relaxed state. Table 17 summarizes the parameters used to optimize the lattice structure, including force field used, quality, summation method and cut-off distance.

<table>
<thead>
<tr>
<th>Geometry optimization parameters</th>
<th>Force field</th>
<th>COMPASS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quality</td>
<td>Ultrafine</td>
<td></td>
</tr>
<tr>
<td>Summation Method</td>
<td>Ewald</td>
<td></td>
</tr>
<tr>
<td>Repulsive cutoff</td>
<td>6Å</td>
<td></td>
</tr>
</tbody>
</table>

Table 17: Geometry optimization parameters
7.3.2 Dynamic simulation parameters

Table 18 shows the dynamic simulation parameters used in this study. Different methods and parameters can be used to run dynamic simulations (NVT, NVE and NPT). In this study, NPT method was used, in which N the number of atoms is, P is the pressure and T is the temperature. The choice of the total time of simulation and the time step is also important in determining whether the simulation has reached equilibrium before calculating average mechanical properties. In accordance with the reduced time of simulation targeted in this study, total simulation time was selected to be between 200 and 500 ps, and the time step equal to 0.5 fs.

<table>
<thead>
<tr>
<th>Dynamic Simulation parameters</th>
<th>NPT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ensemble</td>
<td>NPT</td>
</tr>
<tr>
<td>Temperature</td>
<td>298 K (25°C)</td>
</tr>
<tr>
<td>Pressure (GPa)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Time step (fs)</td>
<td>0.5</td>
</tr>
<tr>
<td>Total simulation time (ps)</td>
<td>200-500</td>
</tr>
<tr>
<td>Thermostat</td>
<td>Nose</td>
</tr>
<tr>
<td>Barostat</td>
<td>Anderson</td>
</tr>
</tbody>
</table>

7.3.3 Results of the molecular simulation

7.3.3.1 Elastic properties of Tobermorite and Jennite with different silicate chains

The results of molecular simulation of the modified chain Tobermorite-like T2 and Jennite-like J2 by COMPASS force field are presented in Table 19. The results reported in chapter 6 about Tobermorite 14A (T∞) and Jennite (J∞) with infinite silica chain, are also presented here to show the effect of the removal of bridging (removing SiO₂) on elastic properties. For comparison, results of the ab-initio method shown in literature are also provided in Table 19.
### Table 19: Elastic properties of Tobermorite and Jennite with different silicate chain lengths

(Appendix IX, X, XI, XII)

<table>
<thead>
<tr>
<th>Mineral</th>
<th>$T_\infty$ (GPa)</th>
<th>$T_2$ (GPa)</th>
<th>$J_\infty$ (GPa)</th>
<th>$J_2$ (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_0$</td>
<td>40.9</td>
<td>42.9</td>
<td>73.3</td>
<td>71.4</td>
</tr>
<tr>
<td>$G$</td>
<td>16.7</td>
<td>12.8</td>
<td>23.9</td>
<td>16.8</td>
</tr>
<tr>
<td>$E$</td>
<td>44.1</td>
<td>34.93</td>
<td>64.7</td>
<td>46.7</td>
</tr>
<tr>
<td>$\nu$</td>
<td>0.32</td>
<td>0.36</td>
<td>0.35</td>
<td>0.39</td>
</tr>
<tr>
<td>Density (g/cm$^3$)</td>
<td>2.24</td>
<td>1.903</td>
<td>2.32</td>
<td>2.048</td>
</tr>
<tr>
<td>C/S</td>
<td>0.83</td>
<td>1.25</td>
<td>1.5</td>
<td>2.25</td>
</tr>
</tbody>
</table>

Ref: $K_0$: Bulk modulus

$G$: shear modulus

$E$: young modulus

$\nu$: poison ratio

As shown in Table 19, the molecular simulation by COMPASS force field used in this study confirms that removing the bridging of silicate tetrahedrals considerably reduced the Young modulus of analog C-S-H minerals. For the infinite to finite silicate chain, the Young modulus of Tobermorite 14A and Jennite was reduced from 44.1 GPa to 34.93 GPa and 64.7 GPa to 46.7, respectively. However, as mentioned above, the experimental Young modulus of C-S-H (LD and HD) measured by Constantinides (2013) ranged between 20 and 30 GPa. These results indicate that even after removing the bridging of silicate tetrahedrals, the Young modulus of Tobermorite $T_2$ and Jennite $J_2$ remained higher than that of C-S-H, but with fewer differences than the infinite silicate chain considered in chapter 5. In addition, the Young modulus of Tobermorite $T_2$ resulting from the mechanical

[a]: (Shahsavari et al., 2009)
[b]: (Manzano et al., 2007)
[c]: (Moon et al., 2015) (experimental and computation)
[d]: (Rejmak et al., 2012)
simulation by COMPASS force field showed a clear agreement with the results of Manzano et al. (2007), who used numerical simulation by GULP code. However, the Young modulus of Jennite J2 resulting from this study (46.7 GPa) shows some discrepancy compared to the value reported (34.7 GPa) from the ab-initio method used by Manzano et al. (2007).

Another notable result concerns the relationship between C/S ratio, silicate chain length and Young modulus. As illustrated in Table 19, as silicate chain length decreases, C/S ratios increases and Young modulus values decrease. For instance, when the silicate chain was modified from infinite to finite and the Young modulus of Tobermorite (T\(\infty\)) and Jennite (J\(\infty\)) were reduced from 44.1GPa to 34.93 GPa and 64.7 GPa to 46.7 GPa, respectively, the C/S ratios increased from 0.83 to 1.25 and 1.5 to 2.25, respectively. More correlations from the table are discussed below.

### 7.3.3.2 Relation between Young modulus and chain length Tobermorite of Jennite

Figure 56 shows the relationship between Young modulus results and silicate chain length of Tobermorite 14A and Jennite-like structures.

![Young modulus vs chain length of Tobermorite and Jennite](image)

**Figure 56: Relationship between Young modulus and silicate chain length of Tobermorite and Jennite**

Figure 56 confirms the remark mentioned above in section 7.3.3.1; there is a clear relationship between Young modulus and chain length. With the removal of the bridging silicate tetrahedral (from infinite to finite structures), the Young modulus of Tobermorite and Jennite was reduced by 20% and 30%, respectively. These results show that the decrease in the chain length has a considerable influence on Young modulus results, especially for Jennite-like structures.
7.3.3.3 Relation between Young modulus density and water

The relationship between Young modulus and density of Tobermorite and Jennite is presented in Figure 57.

![Young modulus vs density](image)

**Figure 57:** Relationship between Young modulus and density of Tobermorite, Tobermorite T2, Jennite and Jennite J2

Figure 57 shows that the decrease in Young modulus (related to the decrease in the chain length of Tobermorite and Jennite) causes a decrease in the density of crystals. For instance, Tobermorite 14A ($T\infty$) density that was 2.24 g.cm$^{-3}$ with infinite silicate chain, was reduced to 1.903 g.cm$^{-3}$ after removing the bridging (T2), which represents a decrement of around 15%. The density of Jennite was reduced from 2.048 g.cm$^{-3}$ for $J\infty$ to 2.32 g.cm$^{-3}$ for J2, which represents a 12% reduction. This result can be explained by the mass loss caused by the removal of the bridging silicate tetrahedral, which also reduced the total density of the crystal. The removal of the bridging silicate chain also caused the water to silicate ratio (W/Si) to increase in the lattice. Excess water can lead to a less dense structure and prevent interaction between atoms, resulting in a structure that is less stiff.

7.3.4 Discussion

In looking at the molecular dynamic simulation by COMPASS force field used in this study, the Young modulus of Tobermorite T2 was very close to the value found by Manzano et al. (2009). However, the Jennite J2 result was remarkably higher than that found by GULP code (Manzano et
There is no clear reason for the similarity with T2 and the discrepancy with J2, especially considering that Manzano et al. used the same simulation by GULP code for both T2 and J2, and in this study, the same dynamic simulation by COMPASS force field was used for both samples.

Removing the silicate chain caused a remarkable reduction in the Young modulus of Tobermorite 14A and Jennite. This conclusion is in line with the literature results. Due to atom mass loss, the removal of the bridging silicate tetrahedral was accompanied by a reduction in the density of Tobermorite 14A and Jennite, which can also reduce the stiffness and Young modulus of the analog C-S-H minerals. The decrements in Young modulus due to the removal of the bridging silicate chain show that the infinite chain provided stability to the structure, and that removal of bridging caused reduced cohesion between the layers of the structure.

Even after removing the bridging silicate tetrahedral, the Young modulus of Tobermorite 14A and Jennite showed higher values than those of C-S-H (LD and HD), which suggests that accounting for porosity for Tobermorite T2 and Jennite J2 after removing the bridging may lead to results closer to those of C-S-H. For this reason, porosity was accounted for in Tobermorite 14A and Jennite with infinite and finite silicate chains.

### 7.4 The accounting of porosity for Tobermorite 14A and Jennite

#### 7.4.1 Mechanical properties of porous materials at microscale level

Concrete is a heterogeneous material composed of random structures at different scales, from macroscopic to nanometer. Micromechanics offers a solution to this heterogeneity; in it, heterogeneous materials are separated into phases and the average of these phases is calculated. This procedure is called homogenization, a mathematical model that averages the properties of composite materials. Porosity is one of the most important properties of concrete. To account for it with micromechanics, two methods can be used: Mori-Tanaka and self-consistent formulas. Although they both rescale bulk and shear modulus using packing factor, the self-consistent method can better
describe C-S-H, according to literature results (Constantinides and Ulm, 2007). For that reason, only
the self-consistent method was used in this study.

Formulas to rescale the elastic modulus are as fellow:

\[
\frac{G_{\text{NEW}}}{G_0} = \frac{1}{2} - \frac{5}{4} (1 - \Theta) - \frac{3}{16} r_0 (2 + \Theta) + \\
\frac{1}{16} \sqrt{144(1 - r_0^2) - 480\Theta + 400\Theta^2 + 408r_0 \Theta - 120r_0 \Theta^2 + 9r_0^2 (2 + \Theta)^2}
\]

\[
\frac{K_{\text{new}}}{K_0} = \frac{4\Theta G}{G_0} + 3(1 - \Theta)r_0
\]

\(\Theta\): Packing factor (%)

\(K_0, G_0\) are the bulk and shear modulus (GPa)

\(r_0 = K_0/G_0\)

\(K_{\text{new}}, G_{\text{new}}\): Bulk and shear modulus After applying the packing factor

After calculating bulk and shear modulus, elastic modulus can be calculated by using equations 3.25
and 3.26, explained previously in chapter 3, section 4.6.

7.4.2 Elastic properties of C-S-H – low density (LD) and high density (HD) C-S-H

Using a nano-indentation tool, Constantinides and Ulm (2006) revealed the existence of four
frequency distributions of Young modulus (indentation modulus) that depend on water to cement
(w/c) ratio. When they performed nano-indentation mapping on the surface area of cement paste
(Figure 58), they found:

a) M between 0 and 13 GPa represents the area with high porosity (macroporosity).

b) M between 13 and 26 GPa represents the LD response.

c) M from 26 to 39 GPa represents the third peaks that characterize high density (HD).

d) M> 39GPa represents the unreacted clinker phases and Portlandite.

Decreased w/c ratio leads to reduction of the low density peak, increase of the high density peak and
a probable increase in the frequency of the clinker peak.
7.4.3 Elastic properties of analog C-S-H minerals accounting for porosity

As explained above, the Young modulus results calculated in the previous section for $T_\infty$, $T_2$, $J_\infty$ and $J_2$ have higher elastic modulus compared to the experimental results of C-S-H. In this section, porosity was included by applying self-consistent formulas for LD ($\Theta = 0.64$) and HD ($\Theta = 0.74$) zones, as mentioned by Constantinides and Ulm (2007) and explained in chapter 1, section 1.10. Table 23 shows the elastic modulus of Tobermorite14A and Jennite with different chain lengths ($T_\infty$, $T_2$, $J_\infty$ and $J_2$). Indentation modulus ($M$) was calculated for comparison with experimental results performed on the cement paste surface. The indentation modulus was calculated using equation 2.2, as explained in chapter 2, section 2.5.7. The relationship between packing factor and porosity is explained in chapter 4, section 3.9, and is defined by the following equation:

$$P = 1 - \Theta$$

In which $P$ is the porosity and $\Theta$ is the packing factor.

For instance, low density (LD) and high density (HD) C-S-Hs have packing factors of $\Theta = 0.64$ and $\Theta = 0.74$, resulting in porosity of $P = 0.36$ and $P = 0.26$, respectively. Low and high density parameters were added to define elastic modulus range compared to the experimental results.
7.4.4 Elastic properties of Tobermorite 14A and Jennite at LD zone

Table 20 presents the modulus of elasticity, indentation modulus and other elastic properties calculated after considering a porosity of 36%.

Table 20: Elastic properties of Tobermorite and Jennite with different chain length at LD zone (P=0.36)

<table>
<thead>
<tr>
<th>Mineral</th>
<th>$T_\infty$</th>
<th>$T_2$</th>
<th>$J_\infty$</th>
<th>$J_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_0$ (GPa)</td>
<td>40.9</td>
<td>42.9</td>
<td>73.3</td>
<td>71.4</td>
</tr>
<tr>
<td>$G_0$ (GPa)</td>
<td>16.7</td>
<td>12.8</td>
<td>23.9</td>
<td>16.8</td>
</tr>
<tr>
<td>$K_{new}$ (GPa)</td>
<td>30.32</td>
<td>35.22</td>
<td>53.41</td>
<td>58.18</td>
</tr>
<tr>
<td>$G_{new}$ (GPa)</td>
<td>4.91</td>
<td>3.84</td>
<td>7.14</td>
<td>5.1</td>
</tr>
<tr>
<td>$E_{new}$ (GPa)</td>
<td>14</td>
<td>11.11</td>
<td>20.51</td>
<td>14.86</td>
</tr>
<tr>
<td>$\nu$</td>
<td>0.423</td>
<td>0.447</td>
<td>0.436</td>
<td>0.457</td>
</tr>
<tr>
<td>$M$ (GPa)</td>
<td>17.02</td>
<td>13.89</td>
<td>25.32</td>
<td>18.80</td>
</tr>
</tbody>
</table>

Stiffness

| LS |
| LS |
| LS |

LS: $13 < M < 26$ GPa (Low stiffness)
HS: $13 < M < 36$ GPa (High stiffness)

Table 20 shows the high impact of the new porosity parameter on the calculation of different elastic properties. By accounting for porosity according to C-S-H LD (porosity = 0.36), the Young modulus of $T_\infty$, $T_2$, $J_\infty$ and $J_2$ decreased by around 69% compared to the simulation that did not account for it. These high decrement percentages reconfirm the importance of density on elastic modulus results.

When considering the targeted Young modulus of C-S-H, it can be noted that regardless of $J_\infty$ results, Young modulus results of Tobermorite 14A and Jennite with different chain lengths ($T_\infty$, $T_2$ and $J_2$) were slightly lower than the reported mean Young modulus of C-S-H, which ranged between 20 and 30 GPa. However, when considering indentation modulus (M) values, which were equal to 17.02 GPa, 13.89 GPa, 25.32 GPa and 18.8 GPa for $T_\infty$, $T_2$, $J_\infty$ and $J_2$, respectively, the mapping of those values...
into the experimental C-S-H phases completed by Constantinides and Ulm (2007) (Figure 59),
showed that all the analog C-S-H minrelas ranged within the low stiffness C-S-H zone (13 < M < 26
GPa). This indicates that all Young modulus values found after accounting for Tobermorite 14A and
Jennite porosity were in the range that characterized the C-S-H LD zone, which confirms that the
objective from accounting the porosity has been reached.

Figure 59: Location of indentation modulus of different Tobermorite 14A and Jennite samples in
experimental C-S-H phases, (P=0.36%).

7.4.5 Elastic properties of Tobermorite 14A and Jennite at HD zone

The elastic properties of Tobermorite 14A and Jennite, calculated with a porosity of 26%, are
presented in Table 21.
Table 21: Elastic properties of Tobermorite and Jennite with different chain length at HD zone (P = 0.26)

<table>
<thead>
<tr>
<th>Mineral</th>
<th>T∞</th>
<th>T 2</th>
<th>J∞</th>
<th>J2</th>
</tr>
</thead>
<tbody>
<tr>
<td>K_o (GPa)</td>
<td>40.9</td>
<td>42.9</td>
<td>73.3</td>
<td>71.4</td>
</tr>
<tr>
<td>G_0 (GPa)</td>
<td>16.7</td>
<td>12.8</td>
<td>23.9</td>
<td>16.8</td>
</tr>
<tr>
<td>Knew(GPa)</td>
<td>33.73</td>
<td>38.23</td>
<td>59.67</td>
<td>63.26</td>
</tr>
<tr>
<td>Gnew(GPa)</td>
<td>8.29</td>
<td>6.44</td>
<td>11.98</td>
<td>8.52</td>
</tr>
<tr>
<td>Enew (GPa)</td>
<td>23.0</td>
<td>18.29</td>
<td>33.68</td>
<td>24.46</td>
</tr>
<tr>
<td>ν</td>
<td>0.386</td>
<td>0.42</td>
<td>0.405</td>
<td>0.435</td>
</tr>
<tr>
<td>M(GPa)</td>
<td>27.02</td>
<td>22.22</td>
<td>40.33</td>
<td>30.19</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Stiffness</th>
<th>HS</th>
<th>LS</th>
<th>&gt;HS</th>
<th>HS</th>
</tr>
</thead>
<tbody>
<tr>
<td>LS:</td>
<td>13 &lt; M &lt; 26 GPa (Low stiffness)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HS:</td>
<td>26 &lt; M &lt; 36 GPa (High stiffness)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 21 shows that in considering the porosity characteristics of C-S-H HD (P=26%), the modulus of elasticity of Tobermorite 14A and Jennite decreased considerably compared to the values from the dynamic simulation mentioned in Table 20 (which did not account for porosity). The decrement percentage was around 48% for all Young modulus results, which is lower than that shown when C-S-H LD was considered (69%). However, it remains a high decrement percentage, especially considering that 26% porosity is the lowest experimental percentage shown for C-S-H (Jennings, 2000).

Table 21 also shows that at low density (p=0.26), regardless of the slightly higher value of J∞, Young modulus results of Tobermorite 14A (T∞, T2) and Jennite (J2) fell in the range of mean C-S-H values. In addition, when inserting indentation modulus (M) values (equal to 27.02 GPa, 22.22 GPa, 40.33 GPa and 30.19 GPa for T∞, T2, J∞ and J2, respectively, in the experimental C-S-H phases completed by Constantinides and Ulm, 2007, Figure 60), T∞ and J2 fell into the high stiffness range, HD C-S-
H and T2 were in the low stiffness range, whereas, J∞ exceeded the range of HS C-S-H. When applying the low porosity of C-S-H (P=0.26) to the elastic properties of analog C-S-H minerals, the Young modulus of infinite and finite silicate chain Tobermorite 14A (T∞ and T2) ranged within C-S-H values at its low and high stiffness state, respectively. Young modulus of Jennite with finite silicate chain (J2) ranged within the C-S-H values, while Jennite with infinite silicate chain did not match with either low stiffness neither high stiffness C-S-H values.

Figure 60: Location of indentation modulus of different Tobermorite 14A and Jennite samples in the experimental C-S-H phases, (P=0.26%).

7.5 Relationship between Young modulus and packing factors

The relationship between Young modulus values and packing factors of LD C-S-H (Θ = 0.64) and HD C-S-H (Θ = 0.74) is presented in Figure 61.

Figure 61: Young modulus vs. packing factor
Based on the correlation presented in Figure 61, it can be observed that as packing factor increased, Young modulus also increased for all analog C-S-H minerals, becoming equal to zero when the packing factor reached a value of 0.5. This result agrees with conclusions made by Constantinides, et al. (2007), who found that this behavior can be justified by the characteristics of the granular material, which becomes unstable when packing density is loosened.

7.6 Discussion

After removing the bridging silicate chain and applying the low and high porosity that characterize C-S-H (measured experimentally), the Young modulus of Tobermorite 14A with infinite and finite silicate chain (T∞ and T2) fell within the range of HS C-S-H and LS C-S-H (high and low stiffness), whereas for Jennite, only the J2-like structure was within the range of HS C-S-H.

Jennite J2 structure with finite silicate chain showed an indentation modulus of 18.80 GPa and 30.19 GPa with high (P=0.36) and low (P=0.26) porosity. Those values indicate that the Young modulus of Jennite J2 falls in the range of LS C-S-H and HS C-S-H when the high and low porosity are considered, respectively. Because the Young modulus of Jennite J2 covered both stiffness ranges and its C/S ratio (2.25) was within the limit of C/S ratios of C-S-H, Jennite J2 structure can be considered the closer mineral to C-S-H (LD C-S-H and HD C-S-H) when porosity if accounted for. However, although J2 matched the high stiffness zone of C-S-H, it could not reach the maximum stiffness of C-S-H defined by its maximum indentation modulus (M=36GPa). Longer silicate chains may be needed to attain extra stiffness such as n=2 (named pentamer, as shown in Figure 13, chapter 2, section 2.5.6). The pentamer chain is composed of five silicate tetrahedrals rather than the two used in this study. This explanation agrees with the results of the experimental investigation performed on C-S-H by Richardson (1999). Indeed, by using the trimethylsilylation (TMS) technique and nuclear magnetic resonance (NMR) spectroscopy, Richardson showed that the dimeric silicate chain (n=1) is the predominate species in C-S-H, formed during the first 24 hours after mixing. However, any C-S-
H formed after that period, and even after six or more months of hydration, consists of mainly linear polymeric species such as pentamer.

Tobermorite T2 with finite silicate chain showed an indentation modulus (M) of 13.89 GPa and 22.22 GPa when applying the low and high porosities that characterize C-S-H. These values fell in the range of 13 and 26 GPa, which represents the low stiffness area of C-S-H. This result indicates that Tobermorite-like structures (T2) cannot reach the high stiffness zone of C-S-H (HS C-S-H). According to Jennings (2000), C-S-H with low density (low stiffness) forms faster during the early hydration age, while high density C-S-H (high stiffness) takes longer to form (potentially over days). This suggests that the T2 resulting from this study only represents the LS C-S-H formed during the early age of hydration.

When applying the low porosity to the Jennite J∞ structure, the Young modulus value was 20.51 GPa, which falls within the low stiffness range of C-S-H. However, at high porosity, the Young modulus of Jennite J∞ was estimated to be 33.68 GPa, which exceeds the high stiffness range. With a minimum porosity $p=0.26$, Jennite J∞ cannot be in the HD C-S-H range. The bridging silicate chain must be removed to reduce the Young modulus.

Tobermorite T∞ with infinite silicate chain length showed a Young modulus of 14 GPa and 23 GPa. When applying low and high porosity on its structure, the Young modulus of Tobermorite T∞ fell within the low and high stiffness ranges of C-S-H. This indicates that Tobermorite T∞ represents the structure of C-S-H in its low and high densities. However, by considering the low C/S ratio of this mineral (equal to 0.83), the structure of Tobermorite 14A with infinite silicate chain cannot represent C-S-H structure, because the C/S ratio of C-S-H, evaluated experimentally, ranges between 1.2 and 2.3.

### 7.7 Summary

Tobermorite 14A and Jennite were found to be the minerals closest to C-S-H in structure. However, their Young modulus values were higher than that of C-S-H. The objective of this chapter was to
reduce the Young modulus of Tobermorite 14A and Jennite using dynamic simulation by COMPASS force field, to be closer to the experimentally measured Young modulus of C-S-H. The reduction of elastic modulus was attempted using two approaches. In the first method, a molecular dynamic simulation with COMPASS force field was run on Tobermorite 14A and Jennite structures with finite silicate chain, in which the bridging silicate chain had been removed. In the second method, the porosity of Tobermorite 14A and Jennite were accounted for.

In the first approach, after omitting the bridging silicate, Tobermorite T2 and Jennite J2 were the new defect structures created from Tobermorite 14A and Jennite. Tobermorite T2 and Jennite J2 are characterized by a dimer silicate chain (n=1). The removal of the bridging silicate chain caused reductions of 20% and 28% in the Young modulus of Tobermorite 14A and Jennite, respectively. The cause of the reduction in Young modulus after omitting the bridging silicate can be attributed to a number of reasons, but reduced density due to mass loss as a result of removing the bridging silicate chain has been suggested as the principal cause. Although removing the bridging silicate chain caused remarkable reductions of Young modulus in Tobermorite 14A and Jennite, results of the molecular dynamic simulation of Tobermorite T2 and Jennite J2 by COMPASS force field (estimated to 34.93 GPa and 46.72GP, respectively), were higher than those in the mean range of C-S-H (LD and HD), measured experimentally to be between 20 and 30 GPa, respectively.

Because of the higher estimated Young modulus of Tobermorite 14A and Jennite, even after removing the bridging silicate, this study focused on the second proposed method, which was to account for porosity in all Tobermorite 14A and Jennite structures with infinite and finite silicate chains (T∞, T2, J∞ and J2). After applying the two types of porosity that characterize C-S-H in its low (LD) (P=0.36) and high density (HD) (P=0.26) states, the Young modulus of Tobermorite T∞, Tobermorite T2 and Jennite J2 fell within the range of LS C-S-H (low stiffness) and HS C-S-H (high stiffness), which satisfy the goals of this study, although the Young modulus of Jennite J∞ still exceeded the range of the high stiffness zone of C-S-H.
Because the C/S ratio of Jennite J2 was in the range of C-S-H and since its Young modulus after accounting for porosity covered the low and high stiffness areas of C-S-H, Jennite J2 is considered to be the closer structure to C-S-H with its LD and HD states.

The structure of Tobermorite T2 (with porosity accounted for) is also considered to be close to C-S-H. However, the Young modulus of Tobermorite T2 cannot reach the HD C-S-H range, which suggests that its structure can represent only the LD C-S-H.

Although Tobermorite T∞ showed a Young modulus within the range of LD and HD C-S-H after accounting for porosity, the C/S ratio of this mineral is lower than that of C-S-H. This suggests that the structure of Tobermorite 14A with infinite silicate chain cannot represent the structure of C-S-H (either LD or HD C-S-H).

The Young modulus of Jennite J∞ exceeded the limit of C-S-H even after accounting for porosity, which indicates that Jennite with infinite silicate chain cannot represent C-S-H unless the bridging silicate chain is removed and the porosity is accounted for.
Chapter 8  General Conclusions

8.1 General conclusions

Concrete or cement paste is viewed only at macro scale; however, the atomic scale controls its engineering properties. Molecular simulation is the tool used to investigate the interaction between atoms. This thesis is focused on the mechanical properties of clinker phases and hydration products such as Portlandite, Ettringite and C-S-H. Because the lack of models of the atomic structure of C-S-H remains an obstacle to concrete optimization, special attention is given to this significant component of cement hydration. An understanding of the nanostructure of C-S-H, the main binding phase and critical hydration product, can help us design proper cement paste formulations. However, due to its amorphous and unstable structure, the nano- and micro-structure of C-S-H has been more frequently studied using experimental techniques such X-ray diffraction (XRD), scanning probe microscopy (SPM) and Nuclear Magnetic Resonance (NMR). To counter its complicated structure, most of the current literature on C-S-H modeling has been drawn from other minerals that are structurally analog to C-S-H structure. Tobermorite family and Jennite are the nano-crystals most commonly used in the structural modeling of C-S-H. The numerical simulation of those minerals can help resolve the problem of C-S-H structure at its nano-scale level. The goal of this thesis was to use the molecular simulation by COMPASS force field in material studio software to numerically study the nanostructure of clinker minerals, Portlandite, Ettringite and, especially C-S-H to determine their elastic properties. The relationship between C/S ratio, packing factor and silicate chain length in C-S-H structure and its modulus of elasticity we also targeted.
Like the numerical methods used in literature to characterize the C-S-H structure, the molecular dynamic simulation by COMPASS force field used in this study was based on the numerical simulation of Tobermorite family and Jennite. The optimization of the geometry was the first step in all the computation work, and thereafter the dynamic simulation was performed in two steps: equilibrium and production. The unit cell and lattice parameters for different clinker phases (C₃S, C₂S, C₃A and C₄AF), hydration products (Portlandite and Ettringite), and analog C-S-H minerals (Tobermorite 9A, 11A and 14A and Jennite) were selected from literature. The final geometry optimization and dynamic simulation parameters were chosen according to adequate simulation time and accurate physical structure. The elastic properties resulting from the dynamic simulation were compared with those of the experimental results mentioned in the literature. The final elastic modulus of the studied elements was calculated by considering the porosity of the cement paste using a special equation.

Based on the numerical results, the following conclusions can be drawn:

1- There was good agreement between the molecular modeling and the experimental results of the elastic modulus of the clinker phases. However, no clear relationship was noticed between C/S ratio and elastic modulus of anhydrate cement.

2- The elastic properties of Ettringite and Portlandite were also in close agreement with the experimental results. The elastic modulus of Portlandite was confirmed to be 2.7 times higher than that of Ettringite.

3- The calculated Young modulus of Tobermorite 9A, Tobermorite 11A, Tobermorite14A, and Jennite were higher than those of C-S-H estimated experimentally. However, the results of this study aligned with the elastic properties mentioned in literature using quantum mechanics principle.

4- Two methods have been tested to reduce the Young modulus of Tobermorite 14A and Jennite to be close to that mentioned experimentally for C-S-H: breaking the silicate chain length
following the rule $3n-1$ and introducing porosity to the structures of Tobermorite 14A and Jennite ($P=0.26$ and $P=0.36$).

5- After removing the bridging, the molecular dynamic simulation of the finite silicate chain Tobermorite_type T2 and Jennite_type J2 by COMPASS force field showed Young modulus values of 34.93 GPa and 46.72GP, respectively. The finite silicate chain resulted in reduced elastic modulus compared to the infinite silicate chain. However, even after removing the bridging silicate tetrahedral, the Young modulus of the analog C-S-H minerals remained higher than those of the experimental results performed on C-S-H, in which the mean value of Young modulus ranged between 20 and 30GPa.

6- Two types of porosity were applied to infinite and finite silicate chain of Tobermorite 14A ($T_\infty$ and T2) and Jennite ($J_\infty$ and J2), which are specific characteristics of C-S-H at its low density (LD) ($P=0.36$) and high density (HD) ($P=0.26$) states. After accounting for the porosity, the calculated Young modulus of Tobermorite $T_\infty$, Tobermorite T2 and Jennite J2 fit within the range of LS C-S-H (low stiffness) and HS C-S-H (High stiffness).

7- The Young modulus of J2 covered both the low stiffness (LD) and high stiffness zones, and its C/S ratio was within the range of C/S ratio of C-S-H. J2 was confirmed to be the analog C-S-H mineral that can most closely represent the structure of C-S-H.

8- The Young modulus of the T2 structure, with porosity accounted for, was the low stiffness range of C-S-H and can therefore only represent the structure of C-S-H at early age hydration.

9- Because the C/S ratio of Tobermorite 14A was low and its Young modulus did not reach the highest stiffness zone, this analog C-S-H mineral cannot represent C-S-H structure.

10- Jennite with infinite chain ($J_\infty$) cannot represent the C-S-H structure because its Young modulus was confirmed to be beyond the range of HS C-S-H (high stiffness zone).

11- Based on the molecular dynamic simulation by COMPASS force field used in this study, one of the most important parameters affecting the Young modulus of Tobermorite 9A, 11A, 14A and Jennite was interlayer spacing, which played an important role in the direction of
compressibility and the stiffness of the mineral. As the interlayer increased, Young modulus decreased. The direction of the compressibility was in the parallel direction of the interlayer.

12- The packing factor, which determines the morphology of C-S-H atoms, also had a considerable effect on Young modulus.

13- Water to silicate and water to calcium ratios were also major parameters that influenced elastic modulus. Excess of water can increase the spacing between the calcium silicate layers and reduce the interaction between atoms. As W/Ca and W/Si decreased in Tobermorite 14A, 11A, 9A and Jennite, the Young modulus increased.

14- Density, which determines the compactness and packing of particles, was also a major parameter affecting the elastic modulus of analog C-S-H minerals. As density increased, the Young modulus increased.

This study confirms the efficiency of the numerical method using COMPASS force field in material studio software in the estimation of the elastic properties of most clinker phases and hydration products. Unlike the different experimental and computational methods, the numerical simulation by COMPASS force field showed good results in reduced time and with minimum staff.

8.2 Recommendations:

For a further understanding of the mechanical performance of different types of C-S-H, more research will be needed into:

- The study of the nano-structure of C-S-H in blended cement paste or Portland cement paste with supplementary cementing materials.

- The effect of some aggressive species such as magnesium and sulfuric acid that can considerably affect the nano-structure of C-S-H.

Considering the different forms of C-S-H with the presence of high amounts of Na, K or Al, it would be prudent to investigate another C-S-H structures like C- (Na, Al)-S-H or C- (Na-K)-S-H.
Appendixes: Input parameters to Materials Studio software and elastic properties

I- Alite (C3S)

I-1- Geometry optimization parameters
1-2- Dynamic simulation parameters (C3S)
I-3- Elastic properties of C3S

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>162.0032</td>
<td>162.5358</td>
<td>162.2695</td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>48.9432</td>
<td>51.1435</td>
<td>50.0433</td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>6.1727</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Young Modulus (GPa)</td>
<td>145.4131</td>
<td>148.4068</td>
<td>122.6092</td>
</tr>
<tr>
<td>Poisson Ratios</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exy</td>
<td>0.2700</td>
<td>0.2755</td>
<td>0.3869</td>
</tr>
<tr>
<td>Eyz</td>
<td>0.4588</td>
<td>0.4137</td>
<td>0.3418</td>
</tr>
<tr>
<td>Exz</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
II- Belite (C2S)

II-1- Geometry optimization parameters of C2S
II-2- Dynamic simulation (C2S)
II-3- Elastic properties of C2S

\[
\begin{align*}
\text{Bulk modulus (GPa)} &= 11.7086 & \text{Voigt} &= 117.4700 & \text{Reuss} &= 64.5893 \\
\text{Shear modulus (GPa)} &= 22.9442 & \text{Poisson Ratios} & \\
\text{Compressibility (1/TPa)} &= 85.4074 & \text{EYx} &= 0.0452 & \text{EYZ} &= 0.0712 \\
\end{align*}
\]

\[
\begin{align*}
\text{Young Modulus (GPa)} &= 9.6849 & \text{EYX} &= 1.9395 & \text{EZx} &= 1.5219 \\
\text{X} &= 415.3420 & \text{EZy} &= 0.2150 & \text{EYz} &= 0.1071 \\
\text{Z} &= 206.8693
\end{align*}
\]
III- Tricalcium Aluminate (C3A)

III-1- Geometry optimization parameters of C3A
III-2- Dynamic simulation parameters of C3A
Appendices

### III-3- Elastic properties of C3A

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss (GPa)</th>
<th>Voigt (GPa)</th>
<th>Hill (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus</td>
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</tr>
<tr>
<td>Shear modulus</td>
<td>63.5656</td>
<td>64.8535</td>
<td>64.2105</td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>5.7045</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Direction</th>
<th>Young Modulus (GPa)</th>
<th>Poisson Ratios</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>187.0596</td>
<td>Exy = 0.2805, Exz = 0.3741</td>
</tr>
<tr>
<td>Y</td>
<td>196.1939</td>
<td>Eyx = 0.2942, Eyz = 0.3125</td>
</tr>
<tr>
<td>Z</td>
<td>183.6364</td>
<td>Ezx = 0.3672, Ezy = 0.2924</td>
</tr>
</tbody>
</table>
Appendices

IV- Aluminoferrite (C4AF)

IV-1- Geometry optimization

![Image of geometry optimization software interface]

![Image of non-bond options interface]

![Image of additional settings in calculation software]

![Image of forcefield parameters interface]
IV-2- Dynamic simulation C4AF
IV-3- Elastic properties of C4AF

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>126.1768</td>
<td>129.4469</td>
<td><strong>127.8119</strong></td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>81.2613</td>
<td>84.6930</td>
<td><strong>82.9772</strong></td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>7.9254</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Young Modulus (GPa)

- X = 210.6131
- Y = 200.4817
- Z = 219.2973

Poisson Ratios

- Exy = 0.1994
- Exz = 0.3470
- Eyz = 0.1898
- Eyx = 0.1184
- Ezx = 0.3613
- Ezy = 0.1295
Appendices

V - Portlandite

V-1- Geometry optimization parameters

V-2- Dynamic simulation (Portlandite)
V-3- Elastic properties of Portlandite

<table>
<thead>
<tr>
<th></th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>20.2641</td>
<td>72.2364</td>
<td>46.2502</td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>11.0004</td>
<td>24.6656</td>
<td>17.8330</td>
</tr>
</tbody>
</table>
VI- Ettringite

VI-1- Geometry optimization
VI-2- Dynamic simulation parameters

![Dynamic simulation parameters](image_url)
VI-3- Elastic properties of Ettringite

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>14.1734</td>
<td>15.8944</td>
<td><strong>15.0339</strong></td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>6.6668</td>
<td>7.9962</td>
<td><strong>7.3315</strong></td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>70.5549</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Young Modulus (GPa)</td>
<td>14.0316</td>
<td>0.3658</td>
<td>0.1220</td>
</tr>
<tr>
<td></td>
<td>15.7726</td>
<td>0.4112</td>
<td>0.1993</td>
</tr>
<tr>
<td></td>
<td>32.5780</td>
<td>0.2834</td>
<td>0.4117</td>
</tr>
</tbody>
</table>
VII- Tobermorite (9A)

VII-1- Geometry optimization

![Geometry optimization setup](image1)

![Forcefield options](image2)

![Non-Bond options](image3)

![Forcefield calculation](image4)
VII-2- Dynamic simulation
VII-3- Elastic properties of Tobermorite 9A

<table>
<thead>
<tr>
<th>Property</th>
<th>Value 1 (GPa)</th>
<th>Value 2 (GPa)</th>
<th>Value 3 (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus</td>
<td>86.5691</td>
<td>88.7962</td>
<td>87.6827</td>
</tr>
<tr>
<td>Shear modulus</td>
<td>28.0707</td>
<td>41.2592</td>
<td>34.6650</td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>11.5515</td>
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<td></td>
</tr>
</tbody>
</table>

Young Modulus (GPa)

<table>
<thead>
<tr>
<th>Axis</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>86.7475</td>
</tr>
<tr>
<td>Y</td>
<td>137.2605</td>
</tr>
<tr>
<td>Z</td>
<td>110.4004</td>
</tr>
</tbody>
</table>

Poisson Ratios

<table>
<thead>
<tr>
<th>Ratio</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exy</td>
<td>0.2952</td>
</tr>
<tr>
<td>Eyx</td>
<td>0.4671</td>
</tr>
<tr>
<td>Exz</td>
<td>0.4504</td>
</tr>
<tr>
<td>Eyz</td>
<td>0.0929</td>
</tr>
<tr>
<td>Ezx</td>
<td>0.3539</td>
</tr>
<tr>
<td>Ezy</td>
<td>0.0747</td>
</tr>
</tbody>
</table>
VIII- Tobermorite 11A

VIII - 1- Geometry optimization

VIII - 2- Dynamic simulation
Appendices

VIII -3- Results: Tobermorite 11A

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>47.3066</td>
<td>49.2933</td>
<td>48.2999</td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>24.7546</td>
<td>26.6847</td>
<td>25.7197</td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>21.1387</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Young Modulus (GPa):  
X = 65.1879
Y = 94.6211
Z = 67.1960

Poisson Ratios:
Exy = 0.2292
EyX = 0.3327
Exz = 0.3209
Eyz = 0.3308
Exy = 0.1313
Eyz = 0.0932
IX- Tobermorite 14A

IX-1- Geometry optimization

IX-2- Dynamic simulation

IX-3- Results (Tobermorite 14A)
X- Jennite

X-1- Geometry optimization
X-2- Dynamic simulation
Appendices

X-3- Results (Jennite)

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>72.3846</td>
<td>74.1932</td>
<td><strong>73.2889</strong></td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>22.3794</td>
<td>25.3921</td>
<td><strong>23.8857</strong></td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>13.8151</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Young Modulus (GPa)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>66.4797</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y</td>
<td>65.1647</td>
<td>0.4221</td>
<td>0.2688</td>
</tr>
<tr>
<td>Z</td>
<td>71.1862</td>
<td>0.4137</td>
<td>0.3205</td>
</tr>
<tr>
<td>Poisson Ratios</td>
<td>Exy = 0.4221</td>
<td>Eyx = 0.4137</td>
<td>Ezx = 0.2878</td>
</tr>
<tr>
<td></td>
<td>Eyz = 0.2688</td>
<td>Ezy = 0.3205</td>
<td></td>
</tr>
</tbody>
</table>

[Image of Forcite Dynamics and Forcite Calculation windows]
XI- Tobermorite 14A (T2 dimer)

XI-1- Geometry optimization
XIV-2- Dynamic simulation (T2)
### XIV-3 - Results (T2)

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>34.7426</td>
<td>51.1061</td>
<td>42.9243</td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>8.6125</td>
<td>17.1444</td>
<td>12.8784</td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>28.7831</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Poisson Ratios

<table>
<thead>
<tr>
<th>Poisson Ratios</th>
<th>Exy</th>
<th>Eyx</th>
<th>Ezx</th>
<th>Eyz</th>
</tr>
</thead>
<tbody>
<tr>
<td>X = 7.8835</td>
<td>0.2683</td>
<td>1.7106</td>
<td>1.2957</td>
<td>-0.4406</td>
</tr>
<tr>
<td>Y = 50.2580</td>
<td>0.2537</td>
<td>1.2152</td>
<td>0.8003</td>
<td>-0.3589</td>
</tr>
<tr>
<td>Z = 23.1822</td>
<td>0.2683</td>
<td>1.7106</td>
<td>1.2957</td>
<td>-0.4406</td>
</tr>
</tbody>
</table>
XV- Jennite modified (J2)

XV-1- Geometry optimization parameters

![Image of Forcite Calculation window with parameters set to Geometry Optimization and Ultra-fine quality]

![Image of Forcite Non-Bond Options window with Electrostatic and van der Waals options showing Ewald summation method and 1.0e-5 kcal/mol Ewald accuracy, 6 Å repulsive cutoff, and 0.5 Å buffer width]

![Image of another Forcite Non-Bond Options window with similar settings as previously described]
XV-2- Dynamic simulation parameters

**Setup**
- **Task:** Dynamics
- **Quality:** Ultrafine
- **Restart:**
  - Restart
  - Append to current trajectory

**Energy**
- **Forcefield:** COMPASS
- **Charges:** Forcefield assigned
- **Quality:** Ultrafine
- **Summation method**
  - Electrostatic: Ewald
  - van der Waals: Ewald

**Dynamics**
- **Ensemble:** NPT
- **Initial velocities:** Current
- **Temperature:** 298.0 K
- **Pressure:** 1.0e-4 GPa
- **Time step:** 0.5 fs
- **Total simulation time:** 200.0 ps
- **Number of steps:** 400000
- **Frame output every:** 5000 steps

**Thermostat**
- **Thermostat:** Nose
- **Temp. difference:** 10.0 K
- **Q ratio:** 0.01
- **Collision ratio:** 1.0
- **Decay constant:** 0.1 ps
### XV-3- Results of J2

<table>
<thead>
<tr>
<th>Property</th>
<th>Reuss</th>
<th>Voigt</th>
<th>Hill</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus (GPa)</td>
<td>66.1729</td>
<td>77.1246</td>
<td>71.6487</td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>9.2584</td>
<td>24.4116</td>
<td>16.8350</td>
</tr>
<tr>
<td>Compressibility (1/TPa)</td>
<td>15.1119</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Young Modulus (GPa)</td>
<td>36.0110</td>
<td>0.6953</td>
<td>-0.0756</td>
</tr>
<tr>
<td>Poisson Ratios</td>
<td>0.6315</td>
<td>0.4643</td>
<td>0.7250</td>
</tr>
<tr>
<td>X</td>
<td>32.7068</td>
<td>Exy</td>
<td>Eyz</td>
</tr>
<tr>
<td>Y</td>
<td>51.0765</td>
<td>Ezx</td>
<td>Ezy</td>
</tr>
</tbody>
</table>
References


Cement Statistics and Information. (n.d.). Retrieved from USGS:
http://minerals.usgs.gov/minerals/pubs/commodity/cement/


Jorge, S., Dolado, a., & Klaas van, B. (2011). Recent advances in modeling for cementitious materials. *Cement and Concrete Research, 41*, 711-726.

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Portland cement, and blends of Portland cement with blast-furnace slag, metakaol. 34(9), 1733-1777. doi:10.1016/j.cemconres.2004.05.034


