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«HIGH PRESSURE AND HIGH TEMPERATURE STUDY OF MAGNESIOCHROMITE AND ITS GEOPHYSICAL IMPLICATIONS»

(Thesis format: Monograph)

by

«N M TAUHID BELAL KHAN»

Department of Earth Sciences «Geophysics»

A thesis submitted in partial fulfillment of the requirements for the degree of « Master of Science »

The School of Graduate and Postdoctoral Studies The University of Western Ontario London, Ontario, Canada

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Abstract

Magnesiochromite (MgCr₂O₄) is commonly found in the Earth's crust, upper mantle, meteorites, and possibly in lunar crust. Synchrotron X-ray diffraction measurements of MgCr₂O₄ using double-sided laser heating diamond anvil cell showed the dissociation of MgCr₂O₄ to Cr₂O₃+MgO at ~15 GPa and to Mg₂Cr₂O₅+Cr₂O₃ below and above ~1500 K, respectively. At above 20 GPa, only a single phase CaTi₂O₄-type structure of MgCr₂O₄ was observed at 1400-2000 K. Fitting the pressure-volume data of CaTi₂O₄ type structure of MgCr₂O₄ with Birch-Murnaghan equation of state yields volume at ambient condition (V₀) = 264.4(8) Å³, bulk modulus (K₀) = 185.4(4) GPa, and pressure derivative of the bulk modulus (K₀') = 4; and for CaTi₂O₄ type structure of natural chromite: V₀ = 261(1) Å³, K₀ = 175.4(2) GPa, K₀' = 4, and mLd-type Mg₂Cr₂O₅ yields: V₀ = 338.9(8) Å³, K₀ = 186.5(6) GPa, K₀' = 4. Obtained phase diagram and bulk modulus data provide important information to better understand the structure and dynamics of the deep earth.

Keywords

DAC, LHDAC, MgCr₂O₄, CaTi₂O₄, modified ludwigite, phase diagram, equation of state, bulk modulus, synchrotron, X-ray, high pressure, high temperature

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List of Abbreviations

ADXD:	Angle dispersive X-ray diffraction
ANL:	Argonne National Laboratory
APS:	Advanced Photon Source
Ar:	Argon
Au:	Gold
Be:	Beryllium
BM:	Bending Magnets
BNL:	Brookhaven National Laboratory
BS:	Beam Splitter
CCD:	Charge-Coupled Device
Chr:	Chromite
CF:	CaFe ₂ O ₄
CT:	CaTi ₂ O ₄
DAC:	Diamond Anvil Cell
D-DIA:	Deformation Diamond
EDM:	Electronic Eischarge Machine
EDXD:	Energy Dispersive X-ray Diffraction
Fp:	Ferropericlase
FWHM:	Full Width at Half Maximum
GPa:	Giga-Pascal
HP:	High Pressure
HT:	High Temperature
ID:	Insertion device
IR:	Infrared
KB:	Kirkpatrick-Baez (a kind of mirror)
LHDAC:	Laser Heating Diamond Anvil Cell
LP:	Low Pressure
LT:	Low Temperature

mLd:	Modified Ludwigite
Ne:	Neon
NF:	Notch filter
NSLS:	National Synchrotron Light Source
PTM:	Pressure transmitting medium
RDA:	Rotation Drickamer Apparatus
rf:	Radio frequency
RHDAC:	Resistive Heating Diamond Anvil Cell
XRD:	X-ray Diffraction

List of Symbols

- c: Speed of light
- E: Energy
- m: Mass
- I: Intensity
- c1/c2: Plank constants
- ε: Emissivity
- λ : Wavelength
- T: Temperature
- d: Spacing of lattice planes
- θ : Angle of specular reflection
- B: Curvature parameter
- P: Pressure
- m: Mass
- M: Mass
- ε: Strain
- f: Eulerian strain
- F: Normalized pressure
- R: Gas constant
- N: Number of moles

- T: Temperature
- K: Bulk modulus
- K_0 ': pressure derivative of the bulk modulus
- V: Volume
- ρ: Density

Chapter 1

1. Introduction

1.1. The Spinel Structure

A spinel mineral is a molecular compound with the formula (A)[B₂]O₄, A and B being metal ions with valences of 2+ and 3+ respectively. Spinel compounds are known to crystallize into a cubic structure with the space group $Fd\overline{3}m$ (O'Neill & Navrostky, 1983; O'Neill & Navrostky, 1984). Two kinds of spinel structures that exist at ambient conditions are normal {(A)[B₂]O₄} and inverse {(B)[AB]O₄} spinels. In the normal spinel A and B ions inhabit the tetrahedral (T) and octahedral (M) sites respectively, while in the inverse spinel half of B ions center the fourfold coordination, leading to relocation of all the A ions to octahedral sites (Hazen, 1999; Levy et al., 2000). In order to completely describe the spinel structure geometry, one has to define the cell parameters and atomic position. An illustration of the spinel structure is presented in Figure 1.1.

Spinels may be synthesized in the laboratory and also occur as minerals. Some of the principal members of the oxide spinel family are: spinel (MgAl₂O₄), gahnite (ZnAl₂O₄), hercynite (FeAl₂O₄), cuprospinel (CuFe₂O₄), magnetite (Fe₃O₄), ulvöspinel (TiFe₂O₄), chromite (FeCr₂O₄), magnesiochromite (MgCr₂O₄), galaxite (MnAl₂O₄), magnesioferrite (MgFe₂O₄), franklinite (ZnFe₂O₄), trevorite (NiFe₂O₄), and the high-pressure silicate ringwoodite [(Mg, Fe)₂SiO₄]. Spinels are usually of normal type, however mutual substitution of A and B cations has been reported in many materials. The extreme cases of cation substitution lead to "inverse" spinels such as CoFe₂O₄, where the Co cation occupies half of the octahedral sites and the Fe cation occupies all the tetrahedral sites. In general most spinels have some degree of inversion. A particular case is NiAl₂O₄, which displays a complete randomization of both cations in octahedral and tetrahedral sites.

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Figure 1.1: The spinel structure. Red spheres are "Oxygen", Gold spheres are "A-atom", and Blue spheres are "B-atom". Solid black line represents the boundary of unit cell. (Structural data from Yamanaka et al., 1983)

Ever since Finger and colleagues (1986) studied the structure of spinel and magnetite at pressure up to 4 GPa, their report attracted much attention and many similar work on high-pressure spinel studies have been performed. The motivation to study these materials comes not only from geophysics, but also from technological applications and fundamental physics. In particular, spinel oxides have been widely studied under pressure because these are constituents of many igneous and metamorphic rocks and because the 660 km seismic discontinuity inside earth's mantle has long been recognized with the decomposition of ringwoodite into (Mg, Fe)SiO₃ perovskite and (Mg, Fe)O magnesiowüstite (Hazen et al., 2000).

This class of structures exists in large quantities in the Earth's transition zone. It plays a significant role in describing the physical properties of our planet (Boehler, 2000). Olivine transforms to wadsleyite at high temperature and high pressure, in the environment of below 410 km deep earth mantle. At higher pressure and temperature environment, corresponding to >520 km deep mantle of earth, wadsleyite transforms into ringwoodite, which has the spinel structure (Chudinovskikh & Boehler, 2001). The formula of olivine, Mg_2SiO_4 , can be rewritten as $SiMg_2O_4$, with Si assuming the role of an A cation and Mg the role of a B cation.

Spinels are also very important in materials science. These are binary oxides, which occur in numerous industrial processes and have important technological applications. Petric and Jacob (1982) reported that spinels are formed as stable corrosion products when alloys are exposed to an oxidizing environment. Spinels provide a protective layer between the alloy and gas, preventing further corrosion. Additionally, owing to their favorable electrical properties, certain spinels have been considered as candidate materials for magneto hydrodynamic electrodes and as promising materials (especially LiMn₂O₄) as cathode for Lithium batteries (Malavasi et al., 2005). Spinels are also used as catalysts for the decomposition of chlorinated organic pollutants (Kim & Ihm, 2001), as magnetic materials (Martinho et al., 2001), super hard materials (Zerr et al., 1999), high temperature ceramics (Kim et al., 2001) and as high pressure sensors (Jahren et al., 1992).

Due to their importance in geosciences, physics, materials science, and crystallography, properties of spinel compounds have been explored extensively. A great number of studies have been performed to find the pressure-induced phase transformations in spinel-type structures (Ringwood & Ried, 1969; Irifune et al., 1991, Catti et al., 1999; Levy et al., 2000; Haavik et al., 2000; Shim et al., 2001; Wang et al., 2002; Ono et al., 2006). Two types of high pressure behaviors have been reported: one involving a pressure-induced phase transformation from spinel structure to high pressure polymorph; and the other involving dissociation to a mixture of constituent oxides. High pressure and high temperature (HP-HT) study of MgAl₂O₄ by Ono et al.

(2006) reveals the transformation of cubic $MgAl_2O_4$ to several new phases at different pressure and temperature conditions (Figure 1.2). Other studies involve determining equations of state for numerous spinels (Irifune et al., 2002; Wang et al., 2003; Speziale et al., 2004) and performing theoretical calculations in aim of describing this wide array of structures (Catti et al., 1999).



Figure.1.2: Phase diagram of MgAl₂O₄ reported by Ono et al. (2006). SP=Spinel-type, MA=MgO+Al₂O₃, CF=CaFe₂O₄-type MgAl₂O₄, EP=ε-type MgAl₂O₄, CT=CaTi₂O₄-type MgAl₂O₄

1.2. Chromite Spinel

Chromites are molecular compounds with the general formula ACr_2O_4 and have a normal type spinel structure (Romeijn, 1953; Miyahara & Ohnishi, 1956). The tetrahedra of oxygen atoms

surround divalent metal ions and the octahedra of oxygen surround chromium ions (Sawaoka et al., 1971). The divalent metals ions can be Fe, Mg, Zn or Mn. The chromite structure is similar to the spinel structure presented in Figure 1.1, where the yellow spheres are chromium ions and gold spheres are magnesium ions.

Chromite spinel [(Mg, Fe, Al)Cr₂O₄] is a common mineral in many meteorites (Rubin, 1997). More importantly, it has a high resistance against weathering & diagenesis and is often the only mineral surviving in fossil meteorites (Thorslund et al., 1984). By studying in the shock veins of the Suizhou meteorite (Figure 1.3), Chen et al. (2003a, 2003b) reported that chromite spinel (56–57 wt.% Cr₂O₃, 29 wt.% FeO, and small amounts of Al₂O₃, MgO, TiO₂, MnO and V₂O₃) transforms to the CaFe₂O₄ (CF) type structure at 12.5 GPa and to the CaTi₂O₄ (CT) type structure above 20 GPa. The Suizhou meteorite is an L6 chondrite with shock classification stage as S3 to S4 (Xie et al., 2001, 2002), which fell in Dayanpo, in the southeast of Suizhou in Hubei, China, on April 15, 1986.



Figure 1.3: High-pressure polymorphs of chromite first reported by Chen et al. (2003), by studying in the shock veins of the Suizhou meteorite

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Yamamoto et al. (2008) found numerous exsolution lamellae of diopsidic clinopyroxene and coesite in podiform chromitites from the Luobusa ophiolite. They were explained by the association of high pressure minerals (diamond, clinopyroxene and coesite) with chromites in the podiform chromitites of the Luobusa ophiolite by the inverse transformation from CT/CF-phase to chromite by a mantle upwelling process (Yang et al., 2007, Yamamoto et al., 2009, and Arai, 2010 & 2013). It is necessary to incorporate SiO₂ and CaO components in the host chromite or its precursor, for occurrence of diopsidic clinopyroxene and coesite exsolution lamellae in chromite. But natural chromite contain only 0.6 wt.%, of SiO₂ and the CaO content is less than several tens of ppm (Arai and Yurimoto, 1994). But MgCr₂O₄-rich CF/CT-phases can contain Ca, Si, Ti, and Fe as CaFe₂O₄ and CaTi₂O₄ solid solutions (Chen et al., 2003a, 2003b). This indicates the possibility of presence of high pressure polymorph of chromite spinel in deep earth mantle.

Spinel was reportedly found in lunar rocks (Figure 1.4), and magnesiochromite spinel is one of the highly plausible candidate (Cloutis et al., 2004, Pieters et al., 2011).



Figure 1.4: Reflectance spectra of spinels (Cloutis et al., 2004). Gray spectra in background are the common lunar minerals.

Chromite is commonly associated with olivine, serpentine, magnetite and corundum. Inclusions of chromite-magnesiochromite (FeCr₂O₄-MgCr₂O₄) solid solutions, in particular, magnesiochromite is among the most common inclusion in diamonds from the cratonic lithosphere (Stachel & Harris, 2008; Griffin & Ryan, 1995; Bulanova, 1995). The chemical analysis of spinels included in diamonds reported by Lenaz et al. (2009) showed that the most abundant component is magnesiochromite (magnesiochromite + chromite reach about 90%) followed by chromite and hercynite (6-7%).



Figure 1.5: Spinels with Cr-numbers of at least about 80 is stable in cratonic Lithosphere (Stachel and Harris 2008)

Chromium bearing spinels [(Mg, Fe)Cr₂O₄] are found in peridotite from the Earth's mantle. It also occurs in layered ultramafic intrusive rocks (Gu and Wills 1988). In addition, it is found in metamorphic rocks such as some serpentinites. Ore deposits of chromite form as early magmatic differentiate. It is commonly associated with olivine, magnetite, serpentine, and corundum. Although typically present as accessory phases, chromium bearing spinels are widely considered to be important petrogenetic indicators (Irvine 1965, 1967; Evans and Frost 1975; Sack and Ghiorso 1991; Bosi et al. 2008), and can target a certain regions in garnet lherzolite for diamond exploration (Griffin and Ryan 1995, Stachel and Harris 2008). Chromium spinels present compositional variations and order-disorder modifications related to the petrologic processes in

which they are involved (Perinelli et al. 2014). Chromium spinels are stable over a wide range of temperatures and pressures in the Earth upper mantle. Besides Cr, the equilibrium intercrystalline exchange of Mg^{2+} and Fe^{2+} cations has been intensely analyzed, and revealed its great potential to be used as geothermometer in rocks containing the paragenesis olivine-spinel (Fabries, 1979; Ballhaus, 1990). Chromium bearing spinels being the only commercial ore of chromium (Duke, 1983) is also important in materials science. Chromium is used in stainless steel, nichrome and chrome plating. Chromites are also used in the production of ferrochrome (an important alloying agent in preparing stainless steel) (Arvanitidis et al., 1996), metallic chromium, chromium chemicals, and refractories (Tathavadkar et al., 2005).

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Due to their importance in geosciences, physics, materials science, and crystallography, properties of chromium bearing spinels have been explored extensively both theoretically and experimentally. Ab initio calculation has been performed by Catti et al. (1999) to examine the phase transformation of chromites. The simulation shows that the chromite can dissociate to constituent oxides at high pressure and room temperature. Wang et al. (2002) first studied the magnesiochromite (MgCr₂O₄) spinel at high pressure with raman and reported that this spinel begins undergoing a phase transformation to CaFe₂O₄(CF) or CaTi₂O₄(CT) structure at 14.2 GPa. The complete transition take place at 30.1 GPa (Figure 1.6).

However, the transformation of magnesiochromite to CF or CT phase was not supported by a more recent study by Yong et al. (2012). Instead, a polymorphic phase transition from cubic structure to tetragonal structure was reported for magnesiochromite at 23.5 GPa and room temperature (Figure 1.7) and at a much lower pressure (12 GPa) for iron-chromite (Kyono et al., 2012).



Figure 1.6: Phase transformation of Magnesiochromite by Raman study (Wang et al., 2002)



Figure 1.7: Phase transition from cubic structure to tetragonal structure of magnesiochromite (Yong et al., 2012)



Figure 1.8: Phase diagram of FeCr₂O₄ at HP-HT. Solid lines represent phase boundaries. Sp = spinel-type FeCr₂O₄; mLd = modified ludwigite-type Fe₂Cr₂O₅; Es = Cr₂O₃ eskolaite; CF = CaFe₂O₄-type FeCr₂O₄; CT = CaTi₂O₄-type FeCr₂O₄. (Ishii et al. 2014)

Another recent synthesis study on magnesiochromite (MgCr₂O₄) by Ishii et al. (2015) showed a different behavior. Magnesiochromite dissociates into a mixture of new modified ludwigite-type Mg₂Cr₂O₅ phase + corundum-type Cr₂O₃ at 1100–1600°C, but it decomposed first into MgO periclase + corundum-type Cr₂O₃ at 1000°C at 12-15 GPa. At about 17–19 GPa, the mixture of Mg₂Cr₂O₅ phase + corundum-type Cr₂O₃ transforms to a CaTi₂O₄ type single MgCr₂O₄ phase (Figure.1.9). Ishii et al. (2015) didn't find any trace of CaFe₂O₄ type structure of MgCr₂O₄.



Figure 1.9: Phase diagram of MgCr₂O₄ at HP-HT. Solid lines represent phase boundaries. A dashed line represents the extrapolated transition boundary of mLd + Es to CT. Sp = spinel-type MgCr₂O₄; mLd = modified ludwigite-type Mg₂Cr₂O₅; Es = corundum type Cr₂O₃ eskolaite; CT = CaTi₂O₄-type MgCr₂O₄; Pc = rocksalt-type MgO periclase. (Ishii et al, 2015)

1.3. Objective of the thesis

The high pressure and high temperature (HP-HT) polymorphs of spinel series could potentially provide a comprehensive pressure gauge and geothermometer system over an extended range considering the presence of the spinel minerals in the earth's mantle and in extraterrestrial bodies, their extensive range of composition, their composition dependence of the transition pressure and temperatue. A systematic and careful HP-HT experimental study of magnesiochromite (MgCr₂O₄) has been performed with double sided laser heating diamond anvil cell incorporated with synchrotron X-ray radiation in order to establish the reliable phase diagram.

A study of the equations of state of high pressure polymorphs of chromite spinel is also essential in geophysics for a better understanding of the mechanical and thermal state of the mantle. So, besides constructing the phase diagram of HP-HT polymorphs of magnesiochromite spinel, we have also determined the P-V equation of state (EOS) of all the traceable HP-HT polymorphs of magnesiochromite and one HP-HT polymorphs of natural chromite. This is the first step to determine the P-V-T equation of state of chromium bearing spinels. This would help with the calculation of density and bulk wave velocity over a wide range of P-T conditions and determination of depth dependence of important thermo-elastic parameters such as thermal expansivity and temperature sensitivity of bulk modulus.

Chapter 2

2. Experimental Method

2.1. Experiments and Instruments



Figure 2.1: Experimental Setup of laser heated diamond anvil cell incorporating with synchrotron XRD.

High pressure and high temperature experiments were performed at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (ANL) and at X17B3, National

Synchrotron Light Source (NSLS) of Brookhaven National Laboratory (BNL) using a diamond anvil cell (DAC). Double-sided laser heating method was employed with synchrotron X-ray diffraction in this study. Laser heating was used to heat the sample inside DAC to synthesize new phase and to study phase transformation at high pressure and high temperature. In this chapter, basic principles of DAC, fundamentals of synchrotron radiation, powder X-ray diffraction and theory of Raman spectroscopy and experimental methods are introduced. Figure 2.1 shows the schematic of the experimental setup.

2.2. The Diamond Anvil Cell

In high pressure science, we can generate pressure using two methods (a) dynamic and (b) static compression. While static compression uses a piston-cylinder mechanism, multi-anvil press and diamond anvil cell (DAC) and dynamic compression uses the shock wave method. The advantages of DAC are high transparency to a wide range of electromagnetic radiation, ability to visually observe the pressure effects and high strength to produce ultra-high pressures above 400 GPa (4 Mbar) (Ruoff et al., 1990), attaining the pressure range found in earth's deep interior.

There are various types of DACs, such as the Mao-Bell type, Merrill-Bassett cell, and custommade symmetric DAC. In this study, symmetric DACs were adopted for different experiments and the geometry is shown in Figure 2.2. To prepare the diamond anvil cell for the experiments, firstly. Two diamonds are mounted onto two tungsten carbide seats corresponding to piston and cylinder of the DAC. Pressure is produced by mechanically tightening four screws to force two diamonds opposing each other thereby compressing the sample between two diamonds. The force required even for the highest attainable pressure is not large due to a small culet size of diamond. The culet size of diamond anvil determines the highest pressure it can attain. Normally, smaller the size higher the attainable pressure is.

Due to their ability to generate high pressures, diamonds are the most important components in a DAC. The rest of the cell serves to align the working tips, maintain this alignment under loading and sustain the load. Thus, it is vital to know the properties of diamonds to select the correct diamonds for various experiments. Generally, the diamonds are classified into two main types.

The type I diamonds contain mainly nitrogen atoms as their impurity, commonly at a concentration of 0.1%. On the other hand, type II diamonds have no measurable nitrogen impurities. A further subdivision categorizes diamonds into type Ia having isolated nitrogen atoms; type Ib having aggregated nitrogen atoms; type IIa being clear diamonds and type IIb having boron impurities. Type I diamonds have two strong absorption regions in infrared (IR) spectra, which are around 2000 cm⁻¹ and 1000-1350 cm⁻¹. Type II diamonds have a clean spectrum below 2000 cm⁻¹ allowing IR measurements of the sample inside DAC. Therefore, type I diamonds are suitable for Raman spectroscopy, while type II diamonds are good for IR spectroscopy. At very high pressures, it has been established that type I diamonds with platelet nitrogen aggregates are more resilient to plastic deformation and therefore can constitute the best anvil material for diamond anvil cells functioning in very high pressure (Mao et al., 1979).



Figure 2.2: (a) Shapes of diamond anvils (b) culet designs (Miletich et al. 2000)

The design of anvil is crucial to determining the maximum pressure a DAC can achieve. Generally, two types of cutting are widely used: the modified brilliant cut and the Drukker cut (Figure 2.2). When compared to a brilliant cut diamond, the Drukker cut diamond has an enlarged table area and an increased anvil angle with the absence of highly stressed shoulders. These modifications can strengthen the diamond anvil and help sustain greater applied loads. Bevelled culets diamonds are usually suitable to work at very high pressures (Mao and Bell, 1978). The advantage of a beveled culet is that the stress concentrations at the edges of the culet are reduced; also the support by the gasket in culet is optimized (Bruno and Dunn, 1984). In this study, type Ia diamonds with Drukker cuts and standard culet were employed for synchrotron X-ray diffraction and Raman spectroscopy.

Another important component of DACs is the gasket. Without the gasket, DAC fails at around 10 GPa (Eremets, 1996). The gasket with a hole in the center serves as a sample chamber, building a lower pressure gradient and supporting the tips of the anvils. Different types of gaskets made of stainless steel, tungsten, rhenium, beryllium, boron-epoxy, etc. are used in different types of experiments. In this study, rhenium gaskets were employed in synchrotron X-ray diffraction experiments because of their high strength properties, while stainless steel gaskets were applied for moderate pressure range experiments conducted in-house Raman spectroscopy.

2.3. Synchrotron X-ray Radiation

When charged particles travel in a curved orbit at relativistic speeds, their tangentially emitted electromagnetic radiation is called 'Synchrotron Radiation'. This radiation covers a wide range of wavelengths with extremely high intensity. It ranges from the infrared to visible and ultraviolet, as well as ranging into the soft and hard X-ray of the electromagnetic spectrum. Ever since its initial discovery at a General Electric (GE) laboratory in 1947 (Elder et al., 1947), synchrotron radiation has become a premier research tool for the study of matter in all varied manifestations, and has been quickly developed from first generation to the fourth. A synchrotron facility (Figure 2.3) is mainly composed of parts described below.

An electron gun generates a large number of free electrons. Electrons are then accelerated by a Linac function to a speed close to the speed of light, followed by a process called injection which involves injecting electrons into a vacuum chamber. Such a vacuum metal chamber is called a booster ring and it circulates the speedy electrons inside a metal tube. The radio frequency cavity system (RF) acts on circulating electrons and restores the energy lost through emission of electromagnetic radiation. The bending magnets force the electrons to circulate in a closed orbit by bending the course of the electrons. The insertion devices, such as wigglers and undulators,

are introduced into straight sections of the storage ring and further modify electron trajectories from straight line thereby inducing emission of further synchrotron radiations. The focusing magnets keep the electrons within a narrow range of the defined path with additional fine tuning of the electron beam trajectory. The beam, which is electromagnetic radiation, exits into the user's experimental chambers.



Figure 2.3: Schematic view of a synchrotron radiation source (Adapted from Mi et al., 2012).

A bending magnet produces a fan of radiation in the horizontal plane, like the sweeping of a search light. Several spatially separated experimental stations can be installed using one bending magnet, each using a different part of the fan of radiation. Wigglers and undulators are periodic

devices, consisting of a series of magnets with alternating polarities, which cause the electrons to curve back and forth. Therefore, the insertion devices can increase the intensity and brightness of the bending magnet radiation by up to 10 orders of magnitude.



Figure 2.4: Comparative illustration of the generation process of synchrotron light by a (a) bending magnet, by insertion devices (b) wiggler and (c) undulator. (Adapted from Mi et al., 2012).

The radiations emitted by successive wiggler magnets do not interfere with each other and the total flux from a wiggler is the sum of emissions from each magnet. Thus, the wiggler generates incoherent radiation with a large angle, allowing several branch lines to operate simultaneously by sharing a small portion of the radiation. However when compared to wiggler magnets, the undulator has a smaller angle of deflection of electron trajectory from the straight line compared

to mc^2/E (m: mass; c: speed of light; E: energy) and the emissions from successive magnets retain coherence and interfere with each other. This interference enhances the intensity in certain regions of wavelengths at the expense of other regions and with the use of sufficient number of magnets. The spectrum becomes concentrated to a single or a few narrow, strong lines. For this reason, radiation from an undulator device can be used by only one station at a time. The flux comparison for bending magnet, wiggler and undulator is shown in Figure 2.4.

Synchrotron radiation provides a very powerful tool to many scientific fields such as physics, biology, chemistry, Earth sciences, material science, biology and medical science. The extraordinary properties of synchrotron radiation include: 1) the extremely high intensity enables a quick and accurate measurement; 2) the high collimation ensures a great resolution in measurement due to its spatial precision. This advantage especially benefits high pressure DAC studies due to small sample sizes; 3) the continuous spectrum makes it possible to conduct experiments with either white beam radiation or monochromatic light. Due to these significant advantages, synchrotron radiation source has been especially suitable for studying materials in small amounts and/or under extreme conditions.

2.4. Laser heating

The sample was heated by using double- sided laser heating facility (Figure 2.5) at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory.

The temperature of the laser heated spot was measured by using the light emitted from the heated spot which was fed into a spectrograph; the resulting intensity-vs.-wavelength curve was then fit to the Planck radiation function:

$$\mathbf{I} = \mathbf{c}_1 \cdot \boldsymbol{\varepsilon} \cdot \boldsymbol{\lambda}^{-5} / (\exp(\mathbf{c}_2 / \boldsymbol{\lambda} \cdot \mathbf{T}) - 1)$$
 (Eq: 2.1)

Where I is intensity, c1 and c2 are constants, ε is emissivity, λ is the wavelength and T is temperature. A graph of $\ln(I \cdot \lambda^5)$ vs λ^{-1} was then constructed, which must be a straight line below ~4000 K if the calibration is good. Temperature of the heating spot is simply the slope of
the curve. The lasers and temperature measurement system should be well aligned with the x-ray beam for temperatures to be correlated with x-ray diffraction patterns. Shen et al. (2001, 2005) and Prakapenka et al. (2008) described in detail about the laser heating system in APS.



Figure 2.5: Experimental setup of laser heating diamond anvil cell in at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (a) Picture (b) Schematic diagram (Prakapenka, 2008)

2.5. Powder X-ray Diffraction

By using a rich array of slits, collimators, focusing mirrors etc, the wide range of synchrotron radiation can be dispersed into X-ray, ultra-violet and infrared light and into different hutches for different types of studies. X-ray is high energy electromagnetic radiation, with wavelengths between roughly 0.1 Å and 100 Å (i.e., in between γ -rays and ultraviolet radiation). Therefore, crystal structures can diffract X-rays due to a similar length scale. X-ray powder diffraction is an

important non-destructive method for determining a range of physical and chemical properties of materials, such as phase identification, cell parameters and crystal structure changes, crystallographic textures, crystalline sizes, macro-stress and micro-strain. When compared to a

crystallographic textures, crystalline sizes, macro-stress and micro-strain. When compared to a conventional laboratory X-ray source (e.g., X-ray tube), synchrotron X-rays are 10¹³ times more brilliant. This high energy radiation is capable of penetrating the pressure container to probe high pressure samples. The interaction between X-rays and electrons of atoms in the samples results in a constructive spectrum when the path difference between two diffracted rays equals an integral multiple of the wavelength of incident X-ray. This selective condition is described by the Bragg equation (Braag & Braag, 1913):

$$n \lambda = 2d \sin \theta$$
 (Eq: 2.2)

where n is the order of diffraction, λ is the wavelength of (elastically) scattered radiation, d is the spacing of lattice planes in a sample and θ is the angle of reflection with respect to these planes. The diffraction angle, defined as the angle between the incident primary beam and the diffracted beam, is therefore equal to 2 θ . Diffraction experiments record scattered X-ray intensity as a function of either 2 θ at a fixed wavelength (ADXD) when monochromatic beam is used or as a function of energy at a fixed 2 θ when a white beam source is used (EDXD). In this study, only ADXD were employed.

Angular dispersive X-ray diffraction (ADXRD) is a very popular non-destructive materials characterization technique that has been used in various fields ranging from engineering to biology. Buras and colleagues (Buras et al., 1977) were the first to use ADXRD techniques with DAC interfaced to synchrotron radiation. Since then synchrotron X-ray diffraction has become a well-established technique and has long been successfully applied in high pressure and high temperature studies in diamond anvil cell (Hu et al., 1993), with numerous synchrotron facilities worldwide providing access to scientists conducting experiments.

In ADXRD (Figure. 2.6), the wavelength of the incident X-ray beam is kept constant by using a monochromator and instead the angle at which the incident photons strike the lattice planes is changed. A single-crystal monochromator is used to select a very small segment of energy for

high resolution ADXRD. Diffraction by a powdered sample produces a number of diffraction rings at various 2 θ angles which are collected by an area detector such as an imaging plate or a charge-coupled device (CCD) leads to a series of interference maxima in the shape of rings, concentric with the incident beam at values of 2 θ that satisfy the Bragg law. By using the Bragg equation, $n\lambda$ =2dsin θ , d-spacings are calculated from the observed θ angles. To reduce the possibility of single crystals like diffraction patterns (spotted), samples should be finely ground. Position of the atoms with the unit cell parameters can be calculated from the relative intensities of the diffraction peaks. With the synchrotron x-ray diffraction combining with laser heating systems it is possible to analyze samples at simultaneously high P-T condition in situ. We have performed the powder ADXRD experiment, double-sided laser-heated assemblage at GSECARS (13 ID-D) of APS.



Figure 2.6: Schematic diagram of angular dispersive X-Ray diffraction technique

High pressure phase identification using synchrotron ADXRD techniques involves many advanced experimental methods, so error evaluation becomes an important issue (Ma et al., 2004). One possible source of error arises when using Bragg's law, which requires careful determination of diffraction angle 2θ to obtain accurate results. Another possible source of error

in a high pressure ADXRD experiment is the substantial heating of DAC which can cause a slight shift in lattice parameter. This occurs due to absorption of much of the X-rays with energy less than 10 keV by diamond anvils and sample (Baublitz et al., 1981).

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2.6. Raman Spectroscopy

When photons of a single wavelength (monochromatic light, in this case a laser) interact with matters, the in-elastically scattering phenomenon is called Raman Effect.

With Raman scattering, the incident photons interact with materials and their energy level is either shifted up or down. The photon interacts with the electron cloud of the functional group bonds, and excites them to a virtual state. Electron then relaxes into an excited vibrational or rotational state, causing the photon to lose parts of its energy and been detected as Stokes Raman scattering. The loss of energy depends on the molecular structure, the functional group, the types of atoms in that molecule and its environment. Therefore, Raman spectroscopy can provide a finger print analysis of vibration or rotation in materials. With a transparent diamond window, materials under extreme pressure in a DAC could be studied by Raman spectroscopy.



Figure 2.7: Layout of the Raman system used in this study for high pressure ruby fluorescence measurements.

In this study, a user-customized Raman system combined with ruby fluorescence method was used to determine ruby pressures in a DAC. The schematic diagram of the Raman system is shown in Figure 2.7. A beam of monochromatic light with a wavelength of 514.532 nm is directed into the sample. Most of the light passes straight through, but some is scattered by the sample. The scattering signals from the sample spot are collected by a set of optics and passed through a spectrometer for analysis. Wavelengths close to the laser line, due to elastic Rayleigh scattering, are filtered out while the rest of the collected signals are dispersed onto a liquid-nitrogen cooled charge-coupled device detector. The instrument control and data collection are performed by the software WinSpec. Pressure determination can be obtained directly from shifts in ruby fluorescence peak.

2.7. Ruby Fluorescence Scale

The use of wavelength shift of ruby R1 fluorescence line to determine pressure was originally developed by Forman, Piermarini, Barnett and Block (Forman et al., 1972). The original calibration of this method involved observing the shift of ruby R1 line with pressure, while simultaneously measuring specific volumes of several metals (Cu, Mo, Pd and Ag) by X-ray diffraction. The absolute pressure was obtained indirectly by making reference to the isothermal equations of state of metals derived from shock-wave data (Mao et al., 1986). After being revised numerous times, the ruby fluorescence method is now widely used as a common pressure scale for high pressures above 10 kbar (1 GPa).

When used in a DAC, the ruby chip is placed inside the sample chamber along with pressure medium and sample, and the fluorescence is excited using a laser or any strong light source. An optical detector collects the signal which is then displayed as a graph showing intensity versus wavelength. The R lines for ruby are quite intense and shift linearly to higher wavelengths under hydrostatic pressure (Jayaraman, 1983). Measurement of the wavelength is commonly achieved by determining the wavelength, λ_0 , of the maximum intensity R₁ line (Munro et al., 1985). This value is then used in conjunction with the equation

$$P = 1904/B \{ [1 + (\lambda - \lambda_0)/\lambda_0]^B - 1 \}$$
(Eq: 2.3)

to determine pressure inside the sample chamber, where B is the curvature parameter (7.665 for hydrostatic, 5 for quasi-hydrostatic), λ is the measured wavelength, and λ_0 is the wavelength of maximum intensity R₁ line at ambient pressure (694.24 nm). Although the ruby fluorescence scale is accurate under hydrostatic conditions, when hydrostaticity diminishes it introduces significant error. When ruby is subjected to non-hydrostatic stress, the R lines become broader. Pure uniaxial stress produces an even greater broadening effect than non-hydrostaticity. This increasing line-width phenomenon makes ruby a useful sensor for detecting the limits of hydrostaticity under pressure (Munro et al., 1985).

2.8. Sample Preparation

2.8.1. Sample

In this study, starting material is magnesiochromite (MgCr₂O₄). Dr. Tony Wither, assistant professor of the department of earth sciences in UWO synthesized the magnesiochromite sample. A stoichiometric mixture of reagent grade MgO and Cr₂O₃ was used to synthesize the magnesiochromite. The chemical mixture was first pulverized with ethanol in an agate mortar and then hard-pressed into a pellet. Synthesized of Magnesiochromite was done by heating the pellet in a flowing atmosphere of pure CO₂ at 1300 °C for 24 h. Sample was inspected by powder X-ray diffraction after quenching in air to ambient conditions in order to confirm the synthesis of magnesiochromite. The achieved lattice parameters for magnesiochromite are a = 8.3332 Å and V = 578.68 Å³, consistent with formerly reported parameters (Lenaz et al., 2004).

2.8.2. DAC

Sample preparation and good sample configuration in DAC are essential to a successful experiment. A symmetrical DAC with 300 μ m flat culet diamonds for higher pressure (>20 GPa), and 500 μ m flat culet diamonds for lower pressure (<20 GPa) was used in the experiment. Before using DAC in this experiment, the culets were aligned with the rotational center of cell to eliminate any stress concentrations which could cause the diamonds to crack under high pressure. This was achieved by first tightening the set screws to a point where the diamonds

could not come in direct contact with each other. Next, the diamonds were aligned using the lateral translation screws. The cylinder was then held stationary and the piston was rotated by 180°. After rotation, the diamond mounted in piston was checked for alignment with the cylinder diamond and if found misaligned, was translated halfway to line up with the center of the stationary cylinder diamond via lateral translation screws. The cell was then flipped over and the cylinder diamond was aligned with diamond in the piston. The piston was held stationary while the cylinder was rotated 180° and if found misaligned, the cylinder diamond was translated halfway to the center of the diamond in the piston (similar to the previous step). The process of aligning the diamonds is iterative in nature, and the above procedure is usually repeated 3-4 times before the diamonds are perfectly aligned with the rotational center of the cell.

After the alignment was complete, small pieces of sandpaper were used to remove any debris and then a q-tip soaked in methanol was used to thoroughly clean the culets. This was repeated several times to ensure that the culets were clean and ready for further procedures. Figure 2.8 shows an unaligned and an aligned DAC.



Figure 2.8: Unaligned and aligned DAC

2.8.3. Gasket Preparation

The introduction of a gasket into the diamond anvil apparatus, which was first demonstrated by Van Valkenburg (1965), was a very important development in the history of DAC; this invention that allows the use of DAC as a quantitative tool in high pressure research. The gasket performs

two functions in a high pressure experiment. With a hole drilled through it provides a chamber for the sample being studied and also provides lateral support to the diamond anvils. This support is given by the gasket extruding from the sample chamber around the side of diamond anvils. This extrusion, acting as a supporting ring, creates stress concentrations towards the sample at the edge of culets thereby preventing any motion of gasket material between anvils and thus prevents failure of anvil (Jayaraman, 1983). Below ~5 GPa, the gasket does not display significant deformation and beyond ~5 GPa, the gasket material yields and undergoes large deformation and flows out of the sample chamber to form a curved feature. The flow rate decreases after ~50 GPa due to bending of the diamond anvils. This process is illustrated in Figure 2.9.



Figure 2.9: Deformation of gasket under pressures during indentation; (a) Before deformation at low pressure; (b) After deformation at high pressure

The first step in preparing the gasket was to create an indentation at its center by subjecting the gasket to a pressure predetermined by the designated maximum pressure in the experiment. This was done by using the DAC in conjunction with a small ruby chip to measure the indentation pressure. After the gasket was indented, it was removed from DAC and a hole was drilled in the center of the indentation with an electrical drilling machine (EDM).

An EDM uses electricity as opposed to mechanical methods to machine metals. In this process, a spark is created between a tool (tungsten bit) and a sample (gasket) of different voltage polarity when they are brought into close proximity. This spark occurs when a tool discharges current to the sample through a dielectric medium, can melt a small portion of the sample provided that the spark contains sufficient energy (Lorenzana, 1994). The process can be applied to both soft and

hard metals makes the EDM is appropriate for drilling high pressure gasket materials such as rhenium, stainless steel and inconel with 2-3 μ m tolerances (Lorenzana, 1994). The picture of an EDM is shown in Figure 2.10.



Figure 2.10: Electronic drilling machine (EDM).

In preparation for drilling a hole with the EDM, the indented gasket was placed on the stage and secured with a small metal mount (positive side of the electric circuit). Then the wire drilling tool (negative side of the electric circuit) was lowered (z-direction) until it almost touched the gasket, and the stage was translated (x and y-directions) until the tool was aligned with the center

of the indentation. Off-center alignment could have caused the sample to migrate under load from the center to the edge of the culet. This phenomenon would limit the highest pressures of the experiment and perhaps caused premature failure of the anvils (Lorenzana, 1994). After centering, a dielectric organic solvent (EDM 185TM by Commonwealth Oil's) was dropped onto the stage and the tool was brought close enough to the gasket to initiate the spark. Figure 2.13 shows the gasket at different stages in the process.

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Figure 2.11: Gasket; (a) Rhenium gasket attached in the middle of steel gasket; (b) after indentation with DAC; (c) After drilling hole

2.8.4. Sample Loading

Chromite sample was ground into fine grains and compressed to form a large flake before placing into the sample chamber. This was done using 1 in³ tungsten carbide blocks. With a predrilled gasket centered on the bottom culet (piston side), the sample flake was placed at the center of gasket hole avoiding contact with the gasket. Similarly, gold flake has been prepared with tungsten carbide block, and placed on the opposite diamond in such a way that it touched the corner of the sample which works as a pressure standard. A small ruby chip was then placed on the culet between the sample and the gasket in order to measure the pressure during Ne gas loading. After placement of the sample and ruby, the cylinder side of the DAC was carefully lowered onto the piston, avoiding any impact of top culet with sample and ruby.



Figure 2.12: Example of sample configuration for the X-ray diffraction study in a DAC.

2.8.5. Pressure medium

The next step in the experimental procedure is to load the pressure transmitting medium into the sample chamber. The pressure medium completely fills the spatial voids in the sample chamber that are left after loading the sample and pressure calibrator. The function of pressure medium is to convert the axial forces applied by squeezing mechanism of DAC into hydrostatic pressure on the sample. It is desirable to have a low shear strength (minimizing deviatoric stresses in the sample chamber) and chemical inertness (non-reactive with the sample). The ideal pressure media are noble gases, which fulfill all these requirements (Boehler, 2000).

In this experiment we used liquefied Neon, which helps achieve hydrostatic conditions up to 60 GPa. The pressures reached throughout the experiment were well within the hydrostatic limits of the pressure medium. When loading the pressure medium, the first step was to loosen the set

screws and slightly tighten the main screws, allowing the culets to come in contact with the gasket but not sample. At this point reference guidelines were marked on each screw, with four matching lines being marked on the DAC. These later allowed the pressure medium inside the sample chamber to be secured without applying any initial pressure. Next, the main screws were loosened, turned by 180° and one set screw tightened until a slight gap was observed between the culets and the gasket. If this gap was too large, the sample may be lost when the pressure medium filled the sample chamber. The set screw was then loosened to allow the pressure medium to be contained inside the sample chamber when the main screws were tightened later on in the procedure. The DAC was then gently placed in a container capable of withstanding cryogenic conditions, while avoiding any impulse forces on the sample. This container was placed inside an insulating shell and a helical copper tube which was connected to an argon tank at one end and entered the container at the other end was inserted between them. The tank valve was opened and neon was allowed to flow into the container for several minutes. Next, a cover was placed over the container to separate the DAC from dry air which could have condensed upon cryogenic cooling and entered the sample chamber. Liquid nitrogen was poured between the two containers submerging the copper tube. At this point the neon began to liquefy due to conductive heat transfer with liquid nitrogen and was allowed to flow until the entire sample chamber was submerged. After submersion was complete, the liquefied neon continued to flow until boiling (caused by the room temperature DAC) stopped. Then the main screws were tightened 180° to their original position, containing the pressure medium inside the sample chamber. Finally, the DAC was removed from container and allowed to reach room temperature.

2.8.6. Determination of Equations of state (EOS)

Equations of state (EOS) of deep planetary materials are essential in understanding observed seismic structure of the Earth in terms of its mineralogy, and geodynamics of the planet's interior. By analyzing the seismic travel times, it is possible to derive the average densities, elastic moduli at the various depths, and pressures of the Earth's interior. In order to build consistent mineralogical models of the Earth interior, those parameters can then be used together with the elastic moduli of mineral assemblages determined from EOS.

The EOS relates the pressure (P), temperature (T) and specific volume (V), which are not all independent and have to be specified in order to define the thermodynamic state of a system. The simplest EOS is the one that describes an ideal gas:

$$PV = nRT \qquad (Eq: 2.4)$$

Where R = gas constant and n = number of moles

The effect of temperature (T) is much less for solids than for gases. The simplest isothermal EOS for solids is given by the definition of the bulk modulus K:

$$K = -\left(\frac{d\ln V}{dP}\right)^{-1} = -\frac{dP}{d\ln V} = -\left(\frac{d\ln M}{dP}\right)^{-1} - \left(\frac{d\ln \rho}{dP}\right)^{-1} = \frac{dP}{d\ln \rho} = -V\left(\frac{dP}{dV}\right)_{T}$$
(Eq: 2.5)

If infinitesimal strains result from the application of hydrostatic pressure on the solid, bulk modulus can be considered as constant. When integrating with $K = K_0$, which means the tangent of the P vs. V curve through the zero point, the EOS becomes:

$$V = V_0 \cdot e^{-(\frac{P}{K_0})}$$
 (Eq: 2.6)

This simple EOS does not consider the increase of the bulk modulus with pressure which limit the validity of this EOS only for low pressures.

2.8.7. Murnaghan integrated linear EOS

Murnaghan proposed an EOS in 1944 that considers the linear dependence of the bulk modulus with pressure. The change of K with P is defined by

$$\mathbf{K}' = \left(\frac{\mathrm{d}\mathbf{K}}{\mathrm{d}\mathbf{P}}\right)_{\mathrm{T}} \tag{Eq: 2.7}$$

K' is the pressure derivative of the bulk modulus and describes the curvature of the P vs. V curve. K' is usually in between 2 and 8 and set to 4, if it cannot be determined with confidence.

A simple formalism is derived when expanding K to first order in P:

$$\mathbf{K} \approx \mathbf{K}_0 + \mathbf{K}'\mathbf{P} \tag{Eq: 2.8}$$

Inserting the K into previous equation and with integration, the Murnaghan integrated linear EOS is obtained:

$$\frac{V}{V_0} = \left(1 + \frac{K'P}{K_0}\right)^{\frac{-1}{K'}}$$
 (Eq: 2.9)

2.8.8. Birch-Murnaghan EOS

Birch-Murnaghan EOS (Birch 1947) is the most commonly used EOS by mineralogists, which has been published in 1947. The displacement of a solid under hydrostatic conditions is not uniform; hence the solid undergoes a strain.

If the properties of solids are considered elastic:

$$\frac{V}{V_0} = (1 - 2\varepsilon)^{\frac{3}{2}}$$
 (Eq: 2.10)

As ε is dilatation which is negative for positive pressures, the compression "f" is introduced:

$$f = -\varepsilon$$
 (Eq: 2.11)
$$f = \frac{1}{2} \left[\left(\frac{V}{V_0} \right)^{\frac{-2}{3}} - 1 \right]$$
 (Eq: 2.12)

The Helmholtz free energy (F) is expanded in powers of f:

$$F = af^{2} + bf^{3} + ..$$
 (Eq: 2.13)
 $P = -\left(\frac{dF}{dV}\right)$ (Eq: 2.14)

An expansion of F to second order provides the second-order Birch-Murnaghan EOS, to third order in the third-order Birch-Murnaghan EOS. Finite strain theory gives a numerical value of
$$K^2$$
 = 4 for the infinitesimal case f \rightarrow 0. Normally, close-packed minerals show experimental values of K² close to 4.

2.8.9. The third-order Birch-Murnaghan EOS

P = 3K₀f(1 + 2f)^{$$\frac{5}{2}$$} $\left[1 + \frac{3}{2}(K' - 4)f\right]$ (Eq: 2.15)

$$P = \frac{3}{2}K_0 \left[\left(\frac{V_0}{V} \right)^{\frac{7}{3}} - \left(\frac{V_0}{V} \right)^{\frac{5}{3}} \right] \left\{ 1 - \frac{3}{4} \left[4 - K' \right] \cdot \left[\left(\frac{V_0}{V} \right)^{\frac{2}{3}} - 1 \right] \right\}$$
(Eq: 2.16)

If K' = 4, the second-order BM EOS is obtained.



2.9. Analysis of X-Ray Diffraction Patterns

Figure 2.13: (a) 2-dimensional XRD pattern collected by CCD, (b) after integration (c) peak position determination using PeakFit program

The 2-dimensional CCD detector collects the pattern of rings of diffracted X-rays of various diameters of the crystals (Figure 2.13). These patterns are then integrated azimuthally using Fit2D software (Hammersley et al., 1996) to better illustrate the relationship between the

diffraction rings and their angle to the main X-ray beam. Each ring appears as a peak on a 1dimensional plot; the relationship between the peak's angle to the main X-ray beam and the crystal structure of the sample is given by the Bragg Diffraction Equation:



Figure 2.14: Bragg's Law reflection. The diffracted X-rays exhibit constructive interference when the distance "abc" is an integer number of wavelengths (λ).

In this equation, λ is the wavelength of the photons, θ is the angle of X-ray diffraction relative to the incident X-ray beam, and d is a measure of the distance between planes in an idealized crystal lattice. Peaks position (θ) is obtained using the PeakFit (Version 4.12), which has been used to calculate the d-spacing using Bragg law. The parameters of the unit cell and the volume have been calculated by least-squares fit using the UnitCell program (Holland and Redfern, 1997). The pressure was determined from the measured d-spacings of Au using the equations of state of Au (Fei et al., 2007).

Chapter 3

3. Phase diagram of MgCr₂O₄

High pressure and high temperature experiments of synthetic magnesiochromite (MgCr₂O₄) were performed at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (ANL). From the X-ray diffraction measurements, we collected the diffraction patterns up to 40 GPa and 2000 K in several heating cycles using double-sided laser heating diamond anvil cell. The goal of the experiments was to synthesize the high pressure and high temperature polymorphs of MgCr₂O₄ and construct the phase diagram.

3.1. Experiment

High pressure and high temperature experiments were performed using double-sided laser heating diamond anvil cells technique at GSECARS laser heating facility (Prakapenka et al., 2008). Synthesized MgCr₂O₄ first ground to fine powder and mixed with ~10 wt.% gold for use as a pressure calibrant. Mixtures of the sample and gold were then loaded into the ~150 μ m hole of a rhenium gasket. Neon gas was also loaded in DAC as a pressure medium with the GSECARS high-pressure gas loading system. The temperature of the laser heated spot was measured from both sides using the light emitted from the heated spot by spectroradiometry. The samples were probed with a monochromatic X-ray beam 3×3 μ m² in cross-section and 0.3344 Å in wavelength, whereas the cross-section of laser heating spot was 20×20 μ m². The X-ray beam was ensured to be in the center of the heating spot by carefully aligned the optical paths using fluorescence generated by the X-rays as they passed through the sample. The 2-dimensional CCD detector used to collect the pattern of rings of diffracted X-rays of various diameters of the crystals. The sample-to-detector distance and orientation of the detector was calibrated using CeO₂ powder.

3.2. Results

XRD data from a representative heating cycle are shown in Figure 3.1. At ambient pressure and temperature, it shows only magnesiochromite structure with peak positions at (111), (220), (311), (222), (400), (331), (422), (333), (440), (531), (533), (622), (444) and (711). When the pressure increased to 23.5 GPa at ambient temperature, the tetragonal phase of chromite appeared. At this high pressure, the sample has been heated until it transforms to a new phase (For miler indices: Appendix A3). Due to the local heating phenomenon of the laser heating technique, the pressure increased a few GPa during heating. The new phase has been confirmed as $CaTi_2O_4$ type structure of MgCr₂O₄.



Figure 3.1: Synthesis of CaTi₂O₄ type structure of MgCr₂O₄



Figure 3.2: $MgCr_2O_4$ first dissociates to Cr_2O_3 +MgO about at 1200 K and 16 GPa, and then to $Mg_2Cr_2O_5$ +Cr_2O_3 about at 1700 K and 16 GPa.

In our first heating cycle (Figure 3.1), we obtained a $CaTi_2O_4$ type structure (CT) due to high initial pressure. We observed the dissociation of MgCr₂O₄ to Cr₂O₃ and MgO (Figure 3.2 & Table 3.1) when the sample was heated at ~1200 K at pressure ~15 GPa. When the temperature was increased to ~1600 K, Cr₂O₃ and MgO joined together in 1:2 ratios to form Mg₂Cr₂O₅. At this stage the phases are Mg₂Cr₂O₅ and Cr₂O₃ (Figure 3.2 & Table 3.1).

Run No.	Pressure (GPa)	Temperature (K)	Phases
MgCrE_024	30.6 (4)	1738 (115)	СТ
MgCrE_126	31.2 (5)	1772 (86)	СТ
MgCrE_127	30.8 (4)	1825 (103)	СТ
MgCrE_128	31.2 (4)	1842 (122)	СТ
MgCrE_129	30.9 (3)	1786 (86)	СТ
MgCrE_130	30.9 (2)	1793 (200)	СТ
MgCrE_131	31.1 (2)	1796 (200)	СТ
MgCrE_157	29.4 (3)	1415 (145)	СТ
MgCrE_158	32.2 (2)	1757 (200)	СТ
MgCrE_177	33.6 (3)	1724 (200)	СТ
MgCrE_178	34.1 (3)	1792 (168)	СТ
MgCrE_179	32.8 (2)	1749 (200)	СТ
MgCrG11	19.2 (2)	1079 (190)	Cr ₂ O ₃ +MgO
MgCrG12	22.7 (3)	1750 (200)	СТ
MgCrG14	18 (2)	1540 (1)	mLd+ Cr_2O_3
MgCrG15	18 (3)	1540 (123)	mLd+ Cr_2O_3
MgCrG16	18.5 (2)	1674 (15)	mLd+ Cr_2O_3
MgCrG17	18.8 (2)	1679 (72)	mLd+ Cr_2O_3
MgCrG27	18.9 (3)	1199 (200)	Cr ₂ O ₃ +MgO
MgCrG28	18 (2)	1412 (195)	Cr ₂ O ₃ +MgO
MgCrG29	17.9 (3)	1133 (200)	Cr ₂ O ₃ +MgO
MgCrG30	17.6 (3)	1109 (192)	Cr ₂ O ₃ +MgO
MgCrG31	16.7 (2)	1099 (200)	Cr ₂ O ₃ +MgO
MgCrG32	15.8 (2)	1442 (91)	Cr ₂ O ₃ +MgO
MgCrG34	15.9 (2)	1702 (135)	mLd+Cr ₂ O ₃
MgCrG35	15.4 (3)	1589 (1)	mLd+Cr ₂ O ₃
MgCrG36	15.6 (2)	1594 (18)	mLd+Cr ₂ O ₃
MgCrG38	16.1 (3)	1729 (200)	$mLd+Cr_2O_3$

Table 3.1: Results of high-pressure high-temperature experiments ($CT = CaTi_2O_4$ type structure, $mLd = MgCr_2O_5$).

MgCrG39	15.4 (2)	1710 (115)	mLd+Cr ₂ O ₃
MgCrG40	15.5 (2)	1701 (36)	mLd+Cr ₂ O ₃
MgCrG41	21.3 (2)	1954 (200)	СТ
MgCrG42	17.8 (3)	1730 (97)	СТ
MgCrG43	19.1 (2)	1748 (133)	СТ
MgCrG57	20.9 (2)	1727 (24)	СТ
MgCrG58	20.3 (3)	1692 (37)	СТ
MgCrG59	21.1 (2)	1616 (45)	СТ
MgCrG60	19.2 (3)	1556 (33)	СТ
MgCrG61	24.8 (2)	1835 (108)	СТ
MgCrG62	24.6 (3)	1862 (37)	СТ
MgCrG63	24.4 (2)	1771 (4)	СТ
MgCrH17	13.1 (3)	1630 (200)	Spinel
MgCrH18	13.2 (2)	1637 (200)	Spinel
MgCrH22	13.3 (2)	1225 (120)	Spinel
MgCrH23	13.5 (3)	1547 (109)	Spinel
MgCrH24	17.9 (2)	1494 (25)	mLd+Cr ₂ O ₃
MgCrH25	17.9 (2)	1385 (54)	mLd+Cr ₂ O ₃
MgCrH27	18.6 (3)	1682 (9)	mLd+Cr2O3

The result has been summarized in table 3.1, which has been used to draw the phase diagram (Figure.3.3) between different high pressure high temperature polymorphs of magnesiochromite. In our first set of experiments the pressure was too high (>30 GPa). After heating the sample at ~1700 K all new phases become pure $CaTi_2O_4$ type structure (CT) of single phase MgCr_2O_4. All following x-ray diffraction experiments have been performed at lower pressures (10 to 20 GPa) and temperatures (1000 to 2000 K). We found the MgO+Cr_2O_3 phases at 16.7 GPa and 1100 K at the run number MgCrG31, and Mg_2Cr_2O_5+Cr_2O_3 phases at 15.4 GPa and 1589 K at the run number MgCrG35. However, at pressure ~20 GPa, the heated sample showed CT at run number MgCrG12, MgCrG41, MgCrG42, MgCrG43, MgCrG57, MgCrG58 and MgCrG59. It is to mention that the magnesiochromite structure is present in the first two cases with MgO+Cr_2O_3 and Mg_2Cr_2O_5+Cr_2O_3, also CaTi_2O_4 type structure with Mg_2Cr_2O_5+Cr_2O_3 close to the boundary. We interpreted that the appearance of new phases at any temperature and pressure are the stable

condition for that new phase. The presence of old phases may be due to the duration of the heating was not long enough. Close to the boundary, the sample takes a longer time to transform completely to new phases due to the higher kinetic energy. When the sample experiences the pressure and temperature condition well above the stable condition, it transformed completely in new phases within few minutes. For example, the transformed phase is always pure $CaTi_2O_4$ type structure of MgCr₂O₄ (Figure 3.1), at higher pressure and temperature (>30 GPa, 1700 K).

In the heating cycle shown in Figure 3.2, the first apperance of the peaks of CT phase in run number MgCrG41 at the condition of pressure 21.3 GPa and temperarure 1953K (table 3.1). But if we take a close look at this heating cycle, Temperature and pressure suddenly dropped in the next run MgCrG42 to 17.8GPa and 1730K. The fomation of the new phase shown smaller volumes and therefore the pressure reduced. There was a sudden flash during the phase transformation in the monitor, which was showing the heating spot on the sample. From the safety concern of diamond anvil cell, the laser power was then lowered down to decrease the temperature. Nevertheless, the peaks of CT phase was still observable in the run number MgCrG42. This is probably due to the kinetic issue of the CT phase. The same phenomenon can be observed in run number MgCrH25 (Appendix 5). Mg₂Cr₂O₅+Cr₂O₃ have already been synthesized at higher pressure and temperature and then temperature dropped in the same heating cycle. So, we excluded those kind of data in which temperature dropped suddenly after synthesization of new phases. We plotted the different phases of MgCr₂O₄ obtained in different pressure and temperature in order to construct the phase diagram (Figure.3.3).

The unit-cell lattice parameters determined from peak positions of cubic MgCr₂O₄, tetragonal MgCr₂O₄, mLd-type Mg₂Cr₂O₅, CT-type MgCr₂O₄, MgO and Cr₂O₃ are summarized in Table 3.2. Obtained unit-cell lattice parameters at ambient condition for mLd-type Mg₂Cr₂O₅ (a_0 =9.62894 Å, b_0 =12.4625 Å and c_0 =2.85644 Å) and CT-type MgCr₂O₄ (a_0 =2.85107 Å, b_0 =9.48930 Å and c_0 =9.67853 Å) is very similar to the cell parameter reported by Ishii et al. (2015) (Table 3.3). In table, numbers in the parenthesis are the errors at the last digit.



Figure 3.3: Phase diagram of MgCr₂O₄. Solid black lines represent the phase boundary from our LHDAC experiment. Dashed red lines represent the phase boundary reported by Ishii et al (2015).

Phase	MgCr ₂ O ₄	MgCr ₂ O ₄	Mg ₂ Cr ₂ O ₅	CaTi ₂ O ₄ type	MgO	Cr ₂ O ₃
Symmetry	Cubic	Tetragonal	Orthorhombic	Orthorhombic	Cubic	Hexagonal
Space group	Fd 3 m	I4 ₁ /amd	Pbam	Cmcm	Fm3m	R 3 c
a ₀ (Å)	8.33324(4)	6.06301(7)	9.62031(4)	2.86465(8)	4.21124(7)	4.96527(8)
b_0 (Å)	8.33324(4)	6.06301(7)	12.387304(7)	9.51504(5)	4.21124(5)	4.96527(6)
c ₀ (Å)	8.33324(4)	6.84422(3)	2.842901(6)	9.69811(6)	4.21124(4)	13.55127(4)

Table 3.2: Lattice parameters, symmetry and space group of different phases obtained from $MgCr_2O_4$ at ambient conditions.

Table 3.3: Lattice parameters, symmetry and space group of different phases reported from references for $MgCr_2O_4$ at ambient conditions. But the Tetragonal $MgCr_2O_4$ is at 22.5GPa.

Dhasa	MaCr.O.	MaCr.O.	Ma.Cr.O.	CaTi ₂ O ₄	MgO	Cr ₂ O ₃
1 Hase	MgCr ₂ O ₄	MgCr ₂ O ₄	$Mg_2CI_2O_5$	type		
Source	Lenaz et	Yong et	Ishii et al.,	Ishii et al.,	Sasaki et	Newham et
	al., (2004)	al., (2012)	(2015)	(2015)	al. (1979)	al. (1962)
Symmetry	Cubic	Tetragonal	Orthorhombic	Orthorhombic	Cubic	Hexagonal
Space	$Fd\overline{3}m$	I41/amd	Pbam	Cmcm	Fm3m	$R\overline{3}c$
group	1 00000	1				
a ₀ (Å)	8.3332 (5)	6.063 (1)	9.62894(7)	2.85107(2)	4.217 (1)	4.7589
b_0 (Å)	8.3332 (5)	6.063 (1)	12.4625(1)	9.48930(8)	4.217 (1)	4.7589
c ₀ (Å)	8.3332 (5)	6.895 (3)	2.85644(5)	9.67853(8)	4.217 (1)	12.991

3.3. Discussion

Dissociation of MgCr₂O₄ to Cr₂O₃ and MgO was observed at \sim 15 GPa and temperature below ~1500 K. Upon heating to higher temperature, which is above ~1500 K, Cr₂O₃ and MgO joined together in 1:2 ratios to form Mg₂Cr₂O₅, which is a new phase. This makes the mixture of Mg₂Cr₂O₅ and Cr₂O₃ as the stable phase at this P-T range. At pressure above 20 GPa, only a single phase, CaTi₂O₄ type structure (CT) was observed at the temperature of 1400-2000 K. Our phase diagram (Figure 3.3) is very similar to that of the phase diagram from Ishii et al (2015), but there are some differences. Our boundary between Cr_2O_3+MgO and $Mg_2Cr_2O_5+Cr_2O_3$ is at slightly higher temperatures and the $CaTi_2O_4$ type structure formed at slightly higher pressures. These discrepancies may be due to the uncertainties of pressure and temperature measurements in high pressure and high temperature study. In LHDAC, we measured the pressure using gold peaks and temperature by measuring light emission from the heated spot. Ishii et al. (2015) performed the synthesis experiment using a multi-anvil press apparatus mentioning that there was no correction done for pressure effect on electromotive force of the thermocouple, which may cause this minor difference. These discrepancies may also result from the kinetic issue in laser heating diamond anvil cell technique (LHDAC). We heated the sample about half an hour for each heating cycles. Because the transformation to the new HP-HT phase takes a longer time close to the boundary, there may need a longer heating at the sample. But the important thing is that the high pressure and high temperature polymorphs yielded from our experiments of MgCr₂O₄ are same polymorphs reported by Ishii et al. (2015), and the boundaries between different polymorphs determined from two independent experiments are also of similar patterns.

Chapter 4

4. Equation of State of CaTi₂O₄ type structure of MgCr₂O₄

High pressure and high temperature experiments of synthetic magnesiochromite (MgCr₂O₄) were performed at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (ANL). In order to determine the equation of State of CaTi₂O₄ type structure (CT) of MgCr₂O₄, CaTi₂O₄ type structure was synthesized at high pressure (~45 GPa) and high temperature (~1800 K). Synthesized CaTi₂O₄ type structure of magnesiochromite was then decompressed from 45 GPa to ambient conditions and a great number of XRD patterns has been collected during the decompression stage (Figure.4.1).

4.1. Experiment

Synthesized MgCr₂O₄ first ground to fine powder and mixed with ~10 wt.% gold for use as a pressure calibrant. Mixtures of the sample and gold were then loaded into the ~150 µm hole of a rhenium gasket. Neon gas was also loaded to use as a pressure medium with the GSECARS high-pressure gas loading system. The samples were probed with a monochromatic X-ray beam 3×3 µm² in cross-section and 0.3344 Å in wavelength, whereas the cross-section of laser heating spot was 20×20 µm². The X-ray beam was ensured to be in the center of the heating spot by carefully aligned the optical paths using fluorescence generated by the X-rays as they passed through the sample. The 2-dimensional CCD detector used to collect the pattern of rings of diffracted X-rays of various diameters of the crystals. The sample-to-detector distance and orientation of the detector was calibrated using CeO₂ powder.



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Figure.4.1: Synthesis of CaTi₂O₄ type structure of magnesiochromite was decompressed from 45 GPa to ambient conditions.

4.2. Results

The cubic MgCr₂O₄ spinel structure transformed completely to an orthorhombic CaTi₂O₄ type structure (*Cmcm*) at ~45 GPa and ~1800 K under hydrostatic condition. The lattice parameters for the CaTi₂O₄ type structure at 45.3(3) GPa are a = 2.69714(3) Å, b = 9.00774(4) Å, and c = 9.15484(8) Å.

The observed peak position, and calculated peak position of $CaTi_2O_4$ type structure of magnesiochromite at ambient conditions has been shown in Appendix A2. The lattice parameters for the $CaTi_2O_4$ type structure of $MgCr_2O_4$ at ambient conditions are $a_0= 2.86464(8)$ Å, $b_0=$

9.51504(5) Å, and c_0 = 9.6981(6) Å. Obtained unit-cell lattice parameters at ambient conditions are slightly higher than those determined by Ishii et al. (2015) (a_0 =2.85107(2) Å, b_0 =9.48930(8) Å and c_0 =9.67853(8) Å), but close enough to confirm the synthesis of CaTi₂O₄ type structure. The crystal structure of CaTi₂O₄-type MgCr₂O₄ has been drawn using XtalDraw with structural refinement data from Ishii et al. (2015) (Figure 4.2). The unit-cell parameters and the volume of synthesised CaTi₂O₄ type MgCr₂O₄ at different pressures are presented in table 4.1. Numbers in the parenthesis are the errors at the last digit.



Figure 4.2: Crystal structure of CaTi₂O₄-type MgCr₂O₄ in b-c planes. Solid silver lines represent the unit cell. Blue spheres are Magnesium, green polyhedra are Chromium, and red spheres are Oxygen. (Structure adapted from Ishii et al., 2015)

P (GPa)	a (Å)	b (Å)	c (Å)	V (Å ³)
0.0001	2.86464 (8)	9.51504 (5)	9.6981 (6)	264.343 (8)
2.3 (2)	2.85109 (7)	9.48029 (2)	9.6627 (5)	261.175 (5)
3.2 (3)	2.84533 (3)	9.46918 (6)	9.64956 (7)	259.988 (6)
3.7 (2)	2.84172 (5)	9.46508 (2)	9.63926 (4)	259.268 (6)
4.1 (2)	2.84002 (4)	9.45863 (3)	9.63453 (4)	258.81 (1)
7.9 (2)	2.82134 (2)	9.41101 (3)	9.5711 (7)	254.129 (7)
10.9 (2)	2.80841 (2)	9.36937 (6)	9.52618 (2)	250.663 (6)
12.9 (3)	2.80038 (7)	9.34292 (7)	9.49741 (6)	248.488 (3)
15.5 (3)	2.79082 (8)	9.31014 (7)	9.46331 (5)	245.884 (8)
17.5 (2)	2.78107 (2)	9.2924 (4)	9.43709 (4)	243.881 (8)
20.0 (2)	2.77295 (2)	9.26504 (5)	9.41066 (2)	241.774 (7)
20.8 (2)	2.76834 (3)	9.25479 (5)	9.40235 (7)	240.892 (7)
22.1 (2)	2.76457 (6)	9.24438 (3)	9.38177 (5)	239.767 (8)
23.6 (2)	2.75812 (4)	9.22226 (6)	9.37078 (6)	238.356 (9)
25.3 (2)	2.75384 (3)	9.20848 (6)	9.3467 (2)	237.02 (6)
27.3 (2)	2.74699 (3)	9.184 (8)	9.32012 (7)	235.131 (2)
29.8 (2)	2.74011 (5)	9.16358 (5)	9.28982 (3)	233.26 (4)
31.6 (2)	2.73261 (5)	9.14099 (4)	9.27891 (6)	231.779 (5)
33.6 (2)	2.72758 (7)	9.12619 (7)	9.24748 (7)	230.192 (6)
34.7 (2)	2.72306 (2)	9.11813 (6)	9.23769 (5)	229.365 (5)
38.7 (2)	2.71211 (7)	9.08071 (7)	9.21051 (3)	226.835 (4)
40.5 (3)	2.7099 (7)	9.05589 (3)	9.19153 (4)	225.565 (6)
43.0 (1)	2.70313 (2)	9.03312 (3)	9.16803 (5)	223.862 (6)
45.3 (3)	2.69714 (3)	9.00774 (4)	9.15484 (8)	222.418 (3)

Table 4.1: Unit-cell parameters of synthesised CaTi₂O₄ type MgCr₂O₄ at different pressure

The change of volume with pressure for $CaTi_2O_4$ type structure of MgCr₂O₄ has been shown in Figure 4.4. Normalized pressure vs. Eulerian strain graphs also shown (Figure 4.3) in order to confirm the value of the pressure derivative of the isothermal bulk modulus K₀'. Normalized pressure vs. eulerian strain gives a horizontal line meaning the K₀' is 4. It is to mention that an upward trend would make it more than 4 and a downward trend makes it less than 4. As the K₀' is 4, we can fit our volume-pressure data with second order Birch-Murnagahan EOS. EOS drawn by Ishii et al. (2015) for CaTi₂O₄ type structure of FeCr₂O₄ also showed in the same figure for comparison. The variation of lattice parameters with pressure of CaTi₂O₄ type structure plotted in Figure 4.5.



Figure 4.3: Normalized pressure vs. Eulerian Strain plot of CaTi₂O₄ type structure of MgCr₂O₄



Figure 4.4: Equation of State of CaTi₂O₄ type structure of MgCr₂O₄. Red circles are P-V data and red line is the second order Birch-Murnagahan EOS fitting. Blue solid line is the EOS drawn by Ishii et al. (2014) for CaTi₂O₄ type structure of FeCr₂O₄



Figure 4.5: Variations of unit-cell lattice parameters with pressures for $CaTi_2O_4$ type structure of $MgCr_2O_4$ (a) a-parameter variation with pressure (b) b-parameter variation with pressure (c) c-parameter variation with pressure (d) variation of unit cell dimensions [red circle (a/a₀), Green rectangle (b/b₀), Blue triangle (c/c₀)] with pressure.

4.3. Discussion

From the Figure 4.3, normalized pressure vs. Eulerian strain showed horizontal relationship. So, we can determine the pressure derivative of the isothermal bulk modulus K_0 ' to be 4. This allows us to fit the volume-pressure data with second order Birch-Murnagahan EOS. The fitting yield $V_0 = 264.4$ (8) Å³, $K_0 = 185.4$ (4) GPa. The fitting curve along with pressure-volume data has been shown in Figure 4.4. Shift to the lower volume and pressure range of our EOS fitting curve then that determined by Ishii et al. (2015) indicates that the volume in each pressure step is lower

for $CaTi_2O_4$ type structure of MgCr₂O₄ than $CaTi_2O_4$ type structure of FeCr₂O₄. The variations of lattice parameters with pressures of $CaTi_2O_4$ type structure plotted in Figure 4.5. All three unit cell parameters yield nicely fitted smooth curves. The decreasing magnitude of the unit cell parameters are slightly different from one another (Figure 4.5.d), suggesting the phase exhibit only weak anisotropy in compression behavior.

Chapter 5

5. Equation of State of modified ludwigite-type (mLd type) Mg₂Cr₂O₅ phase

High pressure high temperature experiments of synthetic magnesiochromite (MgCr₂O₄) were performed at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (ANL). In order to determine the Equation of State of mLd-type Mg₂Cr₂O₅, synthesis experiment has been performed at the stable conditions (P = ~16 GPa and T = ~1500K) of mLd-type Mg₂Cr₂O₅ phase. Synthesized mLd-type Mg₂Cr₂O₅ phase was then decompressed from 16.2 GPa to ambient conditions and a great number of XRD patterns has been collected during the decompression stage (Figure.5.1).

5.1. Experiment

Synthesized MgCr₂O₄ first ground to a fine powder and mixed with ~10 wt.% gold for use as a pressure calibrant. Mixtures of the sample and gold were then loaded into the ~150 µm hole of a rhenium gasket. Neon gas was also loaded to use as a pressure medium with the GSECARS high-pressure gas loading system. The samples were probed with a monochromatic X-ray beam $3\times3 \text{ µm}^2$ in cross-section and 0.3344 Å in wavelength, whereas the cross-section of laser heating spot was $20\times20 \text{ µm}^2$. The X-ray beam was ensured to be in the center of the heating spot by carefully aligned the optical paths using fluorescence generated by the X-rays as they passed through the sample. The 2-dimensional CCD detector used to collect the pattern of rings of diffracted X-rays of various diameters of the crystals. The sample-to-detector distance and orientation of the detector was calibrated using CeO₂ powder.



Figure 5.1: Synthesis of Mg₂Cr₂O₅ and Cr₂O₃ phases was decompressed from 16 GPa to ambient conditions

5.2. Results

From our experiments, the synthesized mLd-type $Mg_2Cr_2O_5+Cr_2O_3$ phase is always mixed with original spinel structure and sometime with $CaTi_2O_4$ type structure. We assume that the presence of original spinel structure was due to the heating duration (Each of the heating cycle was around 1 hour). If we heated the sample long enough the residual phases would disappear. On the other hand, presence of $CaTi_2O_4$ type structure was because of the difficulties of keeping the temperature constant in double-sided laser heating technique. The temperature may vary due to the couplings of laser from both sides. In addition, temperature sometime jumped to very high range abruptly during the increasing of the laser power. At some points during heating, the heat
became too high, subsequently causing increase in pressure resulting in a partial synthesis of $CaTi_2O_4$ type structure in some occasions. The stable pressure range of mLd-type Mg₂Cr₂O₅ is also narrow. Due to the high fluctuation of temperature in laser heating technique, it is difficult to obtain pure phase of mLd-type Mg₂Cr₂O₅. For our purpose the mixed phase is good enough to determine the Equation of State, as long as the cell parameters are determinable for that specific phase at different pressure.



Figure 5.2: Crystal structure of modified ludwigite-type (mLd type) Mg₂Cr₂O₅ phase in a-b plane. Solid silver lines represent the unit cell. Blue spheres are Magnesium, different shades of green spheres are Magnesium or Chromium with different probabilities, and red spheres are Oxygen. (Structure adapted from Ishii et al., 2015)

Observed peak position, and calculated peak position of mLd-type Mg₂Cr₂O₅, Cr₂O₃ and CaTi₂O₄ type structure of magnesiochromite at ambient condition is shown in Appendix A3. The lattice parameters for the mLd-type Mg₂Cr₂O₅ at 16.2(2) GPa are a = 9.38888(7) Å, b = 12.04560(8) Å, and c = 2.77967(5) Å and at ambient conditions are a₀ = 9.6203(4) Å, b₀ = 12.38730(7) Å, and c₀ = 2.84290(6) Å. Obtained cell parameters at ambient condition are slightly lower than that determined by Ishii et al. (2015) (a₀ = 9.62894(7) Å, b₀ = 12.4625(1) Å and c₀ = 2.85644(2) Å). Again, the obtained lattice parameters are close enough with published result by Ishii et al. (2015) to confirm the dissociation to Mg₂Cr₂O₅ + Cr₂O₃. The crystal structure of modified ludwigite-type (mLd type) Mg₂Cr₂O₅ phase has been drawn using XtalDraw (Figure 5.2) using the cell refinement data from Ishii et al., 2015.

The dissociation of MgCr₂O₄ phase gives $Mg_2Cr_2O_5$ and Cr_2O_3 . So, we are also able to determine the EOS of Cr_2O_3 phase in order to compare our EOS with the EOS determined by Dera et al. (2011), which will allow verifying our pressure data. Figure 5.2 shows EOS of Cr_2O_3 phase from those same XRD patterns which are used to determine the EOS of mLd-type $Mg_2Cr_2O_5$ phase.

The variations of volume with pressure for mLd-type $Mg_2Cr_2O_5$ phase have been tabulated in Table 5.1, and the EOS has been presented in Figure 5.5. Normalized pressure vs. eulerian strain graphs also drawn (Figure 5.4) in order to confirm the value of the pressure derivative of the isothermal bulk modulus K_0 '. Normalized pressure vs. eulerian strain gives a horizontal line meaning the K_0 ' is 4. Therefore, we fit our volume-pressure data with second order Birch-Murnagahan EOS. The variations of lattice parameters with pressure of mLd-type $Mg_2Cr_2O_5$ phase plotted in Figure 5.6. The decreasing magnitudes of the unit cell parameters are also presented in Figure 5.6.d.



Figure 5.3: Equation of state of Cr_2O_3 . Red circles are P-V data obtained from our sample, and solid red line is Birch-Murnaghan EOS fittings. Solid green curve is the Equation of State of Cr_2O_3 determined by Dera et al (2011).

Table 5.1: Unit-cell parameters of synthesised MgCr2O5 type MgCr₂O₄ at different pressure

P (GPa)	a (Å)	b (Å)	c (Å)	V (Å ³)
0.0001	9.6203 (4)	12.38730 (7)	2.84290 (6)	338.9 (8)
0.2 (1)	9.61926 (6)	12.37961 (6)	2.84259 (2)	338.5 (5)
0.5 (1)	9.61345 (4)	12.37276 (8)	2.84102 (8)	337.9 (5)
0.8 (2)	9.60564 (5)	12.36719 (5)	2.83991 (8)	337.4 (6)
3.4 (2)	9.56685 (5)	12.30314 (5)	2.83019 (7)	333.1 (4)

4.2 (3)	9.54813 (4)	12.28661 (7)	2.82699 (6)	331.6 (4)
6.2 (2)	9.51204 (8)	12.24082 (5)	2.82027 (5)	328.4 (5)
7.5 (2)	9.49621 (3)	12.21436 (5)	2.81456 (2)	326.5 (7)
8.2 (3)	9.48551 (5)	12.20129 (6)	2.81196 (7)	325.4 (8)
9.5 (2)	9.46453 (6)	12.17683 (7)	2.80648 (8)	323.4 (6)
10.4 (2)	9.45029 (8)	12.16268 (8)	2.80293 (9)	322.2 (7)
12.7 (2)	9.42255 (8)	12.11934 (8)	2.79285 (7)	318.9 (2)
14.3 (3)	9.40663 (2)	12.08241 (3)	2.78731 (8)	316.8 (8)
16.2 (2)	9.38888 (7)	12.04560 (8)	2.77967 (5)	314.4 (4)



Figure 5.4: Normalized pressure vs. Eulerian Strain plot of Mg₂Cr₂O₅ phase



Figure 5.5: Equation of State of $Mg_2Cr_2O_5$ phase



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Figure 5.6. Variations of unit cell parameters with pressure (a) a-parameter variation with pressure (b) b-parameter variation with pressure (c) c-parameter variation with pressure (d) variation of unit cell dimensions [red circle (a/a_0), Green rectangle (b/b_0), Blue triangle (c/c_0)] with pressure.

5.3. Discussion

Figure 5.4 shows EOS of Cr_2O_3 phase from those same XRD patterns, which has been used to determine the EOS of mLd-type $Mg_2Cr_2O_5$ phase. The Birch-Murnaghan EOS fittings of our Cr_2O_3 phase yield very similar result to published EOS of Cr_2O_3 by Dera et al. (2011). That gives a higher confidence level to the EOS of mLd-type $Mg_2Cr_2O_5$ phase.

The change in volume with pressure of mLd-type $Mg_2Cr_2O_5$ has been shown in Figure 5.4. Normalized pressure vs. eulerian strain graphs are also drawn (Figure 5.3) in order to confirm the value of the pressure derivative of the isothermal bulk modulus K_0 '. Normalized pressure vs. eulerian strain gives a horizontal line, meaning the K_0 ' is 4. Second order Birch-Murnagahan EOS fitting yield $V_0 = 338.8$ (8) Å³, $K_0 = 186.5$ (6) GPa. The variation of lattice parameters with pressure of $Mg_2Cr_2O_5$ plotted in Figure 5.6. All three unit-cell lattice parameters yield nicely fitted smooth curve. The decreasing magnitude of the unit cell parameters are slightly different with one another (Figure 5.6.d), suggesting the phase exhibit only weak anisotropy in compression behavior.

Chapter 6

6. Equation of State of CaTi₂O₄ type structure of Natural Chromite

High pressure high temperature experiments of natural magnesiochromite (MgCr₂O₄) were performed at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (ANL). In order to determine the equation of State of CaTi₂O₄ type structure (CT) of natural chromite, CaTi₂O₄ type structure was synthesized at high pressure (~40 GPa) and high temperature (~1800 K). Synthesized CaTi₂O₄ type structure of magnesiochromite was then decompressed from 40 GPa to ambient conditions and numbers of XRD reading has been collected during the decompression stage (Figure.6.1).

6.1. Experiment

The geological location of the natural chromite is Sudbury, which we obtained from Dr. Jordan. Natural chromite first separated from other accessory minerals by picking with needle under microscope. Compositional determination with microprobe showed the presence of Mg, Fe, Al, Cr, and a small amount of Mn and Ti [Mn_{0.01}Mg_{0.52}Fe_{0.61}Ti_{0.01}Al_{0.58}Cr_{1.3}O₄]. Separated chromite then ground to a fine powder and mixed with ~10 wt.% gold for use as a pressure calibrant. Mixtures of the sample and gold were then loaded into the ~150 µm hole of a rhenium gasket. Neon gas was also loaded to use as a pressure medium with the GSECARS high-pressure gas loading system. Two sided laser heating technique was used to heat the sample at high temperatures at GSECARS laser heating facility (Prakapenka et al., 2008). The samples were probed with a monochromatic X-ray beam $3\times3 \mu m^2$ in cross-section and 0.3344 Å in wavelength, whereas the cross-section of laser heating spot was $20\times20 \mu m^2$. The X-ray beam was ensured to be in the center of the heating spot by carefully aligned the optical paths using fluorescence generated by the X-rays as they passed through the sample. The 2-dimensional CCD detector used to collect the pattern of rings of diffracted X-rays of various diameters of the crystals. The sample-to-detector distance and orientation of the detector was calibrated using CeO₂ powder.



Figure 6.1: Synthesis of CaTi₂O₄ type structure of natural chromite was decompressed from 40 GPa to ambient conditions.

6.2. Results

The cubic chromite spinel structure transformed completely to an orthorhombic $CaTi_2O_4$ type structure (*Cmcm*) at ~40 GPa and ~1800 K under hydrostatic condition. The lattice parameters for the $CaTi_2O_4$ type structure at 40.8(2) GPa are a =2.6891(6) Å, b =8.98896(5) Å, and c =9.12122(8) Å.

The observed peak positions, and calculated peak positions of $CaTi_2O_4$ type structure of chromite at ambient condition has been shown in Appendix A2. The lattice parameters for the $CaTi_2O_4$ type structure at ambient conditions are $a_0 = 2.8575(6)$ Å, $b_0 = 9.45065(6)$ Å, and $c_0 =$ 9.666412(7) Å. Obtained cell parameters at ambient condition are very similar than those determined by Ishii et al. (2015) ($a_0 = 2.85107(2)$ Å, $b_0 = 9.48930(8)$ Å and $c_0 = 9.67853(8)$ Å), and slightly lower then the cell parameters obtained for the CaTi₂O₄ type structure of MgCr₂O₄ ($a_0 = 2.86464(8)$ Å, $b_0 = 9.51504(5)$ Å, and $c_0 = 9.6981(6)$ Å) in our previous study.

Table 6.1: Unit-cell parameters of synthesised CaTi₂O₄ type of natural chromite at different pressure

P (GPa)	a (Å)	b (Å)	c (Å)	V (Å ³)
0.0001	2.8575 (6)	9.45065 (6)	9.66412 (7)	261.0 (1)
4.5 (2)	2.8254 (8)	9.40488 (8)	9.58554 (4)	254.7 (2)
5.4 (2)	2.8221 (2)	9.38405 (6)	9.5733 (5)	253.5 (2)
7.9 (2)	2.8111 (6)	9.34343 (2)	9.53145 (8)	250.3 (2)
12.4 (3)	2.7917 (7)	9.28542 (4)	9.45297 (5)	245.1 (4)
13.9 (3)	2.7843 (7)	9.26591 (8)	9.44125 (5)	243.6 (2)
15.3 (2)	2.7778 (5)	9.25539 (4)	9.41098 (4)	241.9 (1)
16.8 (2)	2.7700 (6)	9.23299 (4)	9.3873 (6)	240.3 (2)
20.8 (3)	2.7553 (4)	9.19116 (7)	9.34332 (7)	236.9 (2)
22.1 (2)	2.7482 (4)	9.17713 (5)	9.32668 (8)	235.5 (3)
26.14 (2)	2.7339 (6)	9.14167 (5)	9.27814 (5)	231.8 (3)
26.9 (3)	2.7289 (2)	9.12821 (7)	9.26427 (4)	231.0 (2)
29.1 (3)	2.7231 (5)	9.10749 (4)	9.24125 (5)	229.4 (3)
32.0 (2)	2.713 (8)	9.07676 (8)	9.21015 (6)	227.0 (2)
34.2 (2)	2.7064 (8)	9.05617 (4)	9.18631 (4)	225.4 (3)
36.4 (3)	2.7024 (6)	9.04155 (5)	9.16928 (8)	223.9 (2)
37.5 (2)	2.6979 (5)	9.02689 (4)	9.15931 (5)	223.2 (3)
38.1 (2)	2.6957 (7)	9.00995 (7)	9.1458 (7)	222.5 (2)
39.1 (3)	2.6943 (6)	9.00693 (3)	9.14168 (7)	222.1 (3)
39.9 (3)	2.6912 (4)	8.99229 (4)	9.12996 (4)	221.3 (2)
40.8 (2)	2.6891 (6)	8.98896 (5)	9.12122 (8)	220.9 (2)



Figure 6.2: Equation of State of $CaTi_2O_4$ type structure of natural chromite. Red circles are our P-V data and red line is the second order Birch-Murnagahan EOS fitting. Blue solid line is the EOS drawn by Ishii et al. (2015) for $CaTi_2O_4$ type structure of $FeCr_2O_4$, and green solid line is

the EOS of $CaTi_2O_4$ type structure of $MgCr_2O_4$, which has been presented in chapter 4.



Figure 6.3: Normalized pressure vs. Eulerian Strain plot shows that the derivative of the bulk modulus (K_0) is actually 4.



Figure 6.4. Variation of unit cell parameters with pressure (a) a-parameter variation with pressure (b) b-parameter variation with pressure (c) c-parameter variation with pressure (d) variation of unit cell dimensions [red circle (a/a_0), Green rectangle (b/b_0), Blue triangle (c/c_0)] with pressure.

6.3. Discussion

From the Figure 6.3, normalized pressure vs. eulerian strain showed horizontal relationship. So, we can fix the pressure derivative of the isothermal bulk modulus K_0 ' to 4. This allows us to fit the volume-pressure data with second order Birch-Murnagahan EOS. The fitting yield V_0 = 261.0 (1) Å³, $K_0 = 175.4$ (2) GPa. The fitting curve along with pressure-volume data has been shown in Figure 6.2. Comparing the EOS fitting curve of all three CaTi₂O₄ type structure (Figure 6.2) indicates that the volume in each pressure step is lowest in $CaTi_2O_4$ type structure of natural chromite and highest in CaTi₂O₄ type structure of FeCr₂O₄. Natural chromite contains Al beside Fe and Mg. Therefore the presence of Al in chromite makes it smallest in volume. On the other hand, if the tetrahedral of chromite totally occupied by Fe that yields the largest volume. Though the presence of Mg in the chromite tetrahedral makes the chromite lower in volume than Fe bearing chromite, but the volume is higher than Al bearing chromite. Sean et al. (2011) found the similar relationship of cations and volume of crystal studying the equation of state of postperovskites. The variation of lattice parameters with pressure of $CaTi_2O_4$ type structure plotted in Figure 6.4. All three unit cell parameters yield nicely fitted smooth curve. The decreasing magnitude of the unit cell parameters are slightly different from one another (Figure 6.4.d), suggesting the phase exhibit only weak anisotropy in compression behavior.

Chapter 7

7. Geophysical implications

Researchers (Yang et al., 2007, Yamamoto et al., 2009, and Arai, 2010 & 2013) explained the association of high pressure minerals (diamond, clinopyroxene and coesite) with chromite in the podiform chromitites of the Luobusa ophiolite by the inverse transformation from CT/CF-phase to chromite by mantle upwelling process. They explained that the SiO₂ and CaO components are necessary to be incorporated in the host chromite or its precursor for occurrence of diopsidic clinopyroxene and coesite exsolution lamellae in the chromite. But natural chromite contain only 0.6 wt.%, of SiO₂ and the CaO content is less than several tens of ppm (Arai and Yurimoto, 1994). According to Ishii et al. (2014, 2015), CT/CF-phase can't revert back to a cubic chromite structure during mantle upwelling process. So, they concluded that the formation pressure for the Luobusa ophiolite chromities is not more than 12-15 GPa (360–450 km deep). But our experiments indicate otherwise.

After synthesis of a CaTi₂O₄ type structure, we heated the sample at lower pressure (Appendix A2). It showed a growing phase of Cr₂O₃ with time. But we didn't manage to find neither the MgO nor Mg₂Cr₂O₅ phase, which are the associated phases with Cr₂O₃ for two different kind of dissociation identified beforehand. So, we suspect the other phase is MgO, which is not identifiable due to the overlapping with the peaks of CaTi₂O₄ type structure. This is an obvious indication of the dissociation back from CT-phase to Cr₂O₃+MgO. If we look very closely at Figure 3.2 and Appendix 1, we can see that the both kind of dissociates phases Cr₂O₃+MgO and Mg₂Cr₂O₅+Cr₂O₃ can move from one stable condition to another. In Figure 3.2, the dissociate phases are Cr₂O₃+MgO at the beginning of heating moving to Mg₂Cr₂O₅+Cr₂O₃ with increasing temperature. In contrast, Mg₂Cr₂O₅+Cr₂O₃ appeared first at high temperature but peak intensity of Mg₂Cr₂O₅ diminishes with increasing intensity of Cr₂O₃ with time during the heating at lower temperatures (Appendix 1). These indicate that the HP-HT polymorphs of magnesiochromite are very likely able to revert back to low pressure - low temperature (LP-LT) phases, if given

enough time. The chemical composition of natural chromite in the Luobusa ophiolite chromitites reported by Yamamoto et al. (2009) is $(Mg_{0.77},Fe^{2+}_{0.22})_{0.99}(Al_{0.42},Cr_{1.46},Fe^{3+}_{0.11})_{1.99}O_4$, which can be approximated as a solid solution of the MgCr₂O₄-FeCr₂O₄-MgAl₂O₄ system. So, it is very likely that the HP-HT polymorphs of other spinel group members would show similar behavior. Mantle upwelling is a very slow process, which gives the HP-HT polymorphs of spinel enough time to revert back to LP-LT phases. Therefore, our interpretation is that the formation pressure for the Luobusa ophiolite chromitites is very likely more than 12-15 GPa (360–450 km deep).

We showed in chapter 3, MgCr₂O₄ dissociate to Cr₂O₃+MgO and Mg₂Cr₂O₅+Cr₂O₃ phases. In contrast FeCr₂O₄ doesn't show Cr₂O₃+MgO dissociation (Ishii et al. 2014). FeCr₂O₄ dissociates into Fe₂Cr₂O₅+Cr₂O₃ at about 12-16 GPa. At 17–18 GPa, the two dissociate phases combine into CaFe₂O₄-type below 1600 K and CaTi₂O₄-type above 1600 K. The MgAl₂O₄ spinel also has been studied at high pressures and high temperatures by Ono et al. (2006). MgAl₂O₄ showed two different decomposed phases MgO+Al₂O₃ at pressure above 15 GPa and Mg₂Al₂O₅ + Al₂O₃ at about 20 GPa and 2400 K. Those decomposed phases joined together to form a single phase $MgAl_2O_4$ of CaFe₂O₄ type structure at around 30 GPa and then transformed to CaTi₂O₄ type structure above 45 GPa. From the above discussion, it is clear that all these three spinels behave slightly differently at high pressure and high temperature (Appendix A8). Only the common HP-HT polymorph in these three different spinel groups is dissociation to mLd type structure $(Mg_2Al_2O_5/Mg_2Cr_2O_5/Fe_2Cr_2O_5) + oxides (Al_2O_3/Cr_2O_3)$. When Mg is present in tetrahedra (MgAl₂O₄ and MgCr₂O₄), it showed the dissociation into two different oxides (MgO+Cr₂O₃ or MgO+Al₂O₃). FeCr₂O₄ doesn't show that kind of dissociation behavior. So, we may logically conclude that Mg is playing the main role for this kind of dissociation. MgCr₂O₄ doesn't show CaFe₂O₄ type structure at high pressure. The absence of either Fe in tetrahedra or Al in octahedral site may not allow the formation of a CaFe₂O₄ type structure. Both of these structural changes in spinel might be due to the ionic radius of different cations (Appendix A7). Fe²⁺ in tetrahedral normally is in high spin state at ambient condition and ionic radius 92 pm, which is bigger than the ionic radius of Mg²⁺ (86 pm). But at HP-HT iron can shift to low spin state (Yamanaka et al., 2008; Lin et al., 2005), when the ionic radius become only 75 pm. It makes the Mg²⁺ largest cation in size at HP-HT condition in spinel group minerals, which may cause the spinel destabilize and dissociates to the constitute oxides. Similarly, Fe^{2+} and Al^{3+} (67.5 pm) is

smaller in size than Mg^{2+} and Cr^{3+} (75.5 pm), respectively. This may give the spinel enough room to transform to CaFe₂O₄ type structure. This explanation can be verified studying the natural spinel samples at high pressure and high temperature, which is a solid solution of different spinel phases {(Fe,Mg)(Al,Cr)₂O₄}.

Comparing the Birch-Murnagahan EOS fitting curve of all three $CaTi_2O_4$ type structure in Figure 6.2 indicates that the volume in each pressure step is highest in FeCr₂O₄ and lowest in natural chromite. Though the presence of Mg in the chromite tetrahedral makes the chromite lower in volume than Fe bearing chromite, but the volume is higher than Al bearing natural chromite. Therefore the presence of Al in chromite also makes it smallest in volume. On the other hand, EOS fitting showed that the CaTi₂O₄ type structure of MgCr₂O₄ {K₀ = 185.4 (4) GPa} has the lower bulk modulus value then the CaTi₂O₄ type structure of FeCr₂O₄ (K₀ = 199 GPa) and CaTi₂O₄ type structure of natural chromite {K₀ = 175.4 (2) GPa}. Compressibility is the opposite of bulk modulus. Hence, it may conclude that the presence of Fe in tetrahedra makes the spinel less compressible. Natural chromite contains Al, which indicate that the presence of Al makes the spinel highly compressible in nature.

From the discussion above, it is highly likely that HP-HT polymorphs of chromite bearing spinel are present in the deep mantle. This makes it geologically very important to determine the phase diagram and equation of state of these high pressure Cr-bearing polymorphs. The experimentally calibrated high pressure polymorphs of chromite series minerals could be an ideal pressure gauge for shock-metamorphosed terrestrial rocks and meteorites and also for mantle rocks covering the important pressure range throughout the transition zone. Determination of the isothermal bulk modulus is the first step to determine the other thermodynamics parameter of high pressure polymorphs of magnesiochromite in this study will be of great help to facilitate the determination of the speed of sound and other mechanical waves in the high pressure form of magnesiochromite.

Chapter 8

8. Conclusion

High pressure high temperature experiments of synthetic magnesiochromite (MgCr₂O₄) were performed up to 40 GPa and 2000 K at GSECARS, 13-ID-D, Advanced Photon Source (APS) of Argonne National Laboratory (ANL) and at X17B3, National Synchrotron Light Source (NSLS) of Brookhaven National Laboratory (BNL) using double sided laser heating diamond anvil cell (LHDAC).

In the HP-HT experiment of MgCr₂O₄, we observed the dissociation to $Cr_2O_3 + MgO$ at ~15 GPa and temperature below 1440 K, and Mg₂Cr₂O₅ + Cr₂O₃ when the sample heated above 1600 K. At pressure above 20 GPa, only a single phase, CaTi₂O₄ type structure (CT) was observed at the temperature of 1400-2000 K. Our phase diagram is very similar to that of the phase diagram from Ishii et al (2015) with a minor discrepency. Our boundary between Cr_2O_3+MgO and $Mg_2Cr_2O_5+Cr_2O_3$ is at slightly high temperature, and the synthesis pressure of $CaTi_2O_4$ type structure was also slightly higher.

The lattice parameters for the CaTi₂O₄ type structure at ambient conditions are $a_0 = 2.86464(8)$ Å, $b_0 = 9.51504(5)$ Å, and $c_0 = 9.6981(6)$ Å; CaTi₂O₄ type structure of natural chromite are $a_0 = 2.8575(6)$ Å, $b_0 = 9.45065(6)$ Å, and $c_0 = 9.666412(7)$ Å.; and mLd-type Mg₂Cr₂O₅ are $a_0 = 9.6203(4)$ Å, $b_0 = 12.38730(7)$ Å, and $c_0 = 2.84290(6)$ Å. Both of the results are similar to that of the lattice parameter reported by Ishii et al. (2015). Fitting the P-V data with Birch-Murnagahan EOS of CaTi₂O₄ type structure of MgCr₂O₄ yields $V_0 = 264.4(8)$ Å³, $K_0 = 185.4(4)$ GPa, K_0 ' = 4; and of CaTi₂O₄ type structure of natural chromite yields $V_0 = 261(1)$ Å³, $K_0 = 175.4(2)$ GPa, K_0 ' = 4. We have also determined the EOS parameter for mLd-type Mg₂Cr₂O₅ phase $V_0 = 338.8(8)$ Å³, $K_0 = 186.5(6)$ GPa, K_0 ' = 4. All of the phases showed weak anisotropy in compression behavior. It has been also interpreted that the CT phase of magnesiochromite may revert back to the lower pressure lower temperature phases; support the presence of high pressure high temperature polymorphs of chromite spinel in deep mantle. Chromite spinel [(Mg, Fe, Al)Cr₂O₄] is a common mineral in many meteorites and mantle rocks. It is also a possible candidate in the lunar crust. High pressure polymorphs of chromite have been observed in a shocked meteorite and it is very likely that those polymorphs of chromite also exist in the earth's mantle. Obtained phase diagram and bulk modulus will be of great help to understand the physical properties of mantle and shock-metamorphosed terrestrial rocks. This data also possibly have implication for the lunar deep interior.

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Appendices







Appendix A2: Cr_2O_3 peaks showed up after reheating the synthesized $CaTi_2O_4$ type structure of $MgCr_2O_4$

d(obs)	hkl	d(calc)	res(d)
4.84539	002	4.84905	00366
4.27304	021	4.27126	0.00179
3.39788	022	3.39597	0.00191
2.74008	110	2.74303	-0.00295
2.67359	023	2.67384	-0.0002
2.63722	111	2.63935	-0.00212
2.38076	040	2.37876	0.00201
2.16306	024	2.16019	0.00288
2.12425	130	2.12589	-0.0016
2.08864	113	2.09134	-0.00270
2.0749	131	2.07652	-0.002
1.94639	132	1.94689	-5E-04
1.81662	114	1.81644	0.00017
1.79706	025	1.79609	0.00097
1.77648	133	1.77611	0.00037
1.6169	006	1.61635	0.00055
1.58458	150	1.5851	-0.0006
1.5641	151	1.56434	-0.0002
1.50657	152	1.50661	-5E-05
1.50362	045	1.50324	0.00039
1.43212	135	1.43275	-0.0006
1.42431	063	1.42375	0.0006
1.39313	116	1.3925	0.0007
1.37754	202	1.37357	0.00397
1.33739	046	1.33692	0.00047
1.3262	154	1.32667	-5E-04
1.28318	136	1.28659	-0.00341
1.26116	223	1.26249	-0.0013

Appendix A3: Calculated and observed peak positions of synthesised CaTi₂O₄ type MgCr₂O₄.

1.22699	240	1.22705	-6E-05
1.21637	241	1.21732	-1E-03
1.19554	047	1.19719	-0.002
1.19011	080	1.18938	0.0007
1.18075	081	1.18053	0.0002
1.16105	137	1.16063	0.0004
1.12054	225	1.11974	0.0008
1.10994	118	1.10873	0.0012

Appendix A4: Calculated and observed peak positions of synthesised $Mg_2Cr_2O_5+Cr_2O_3$ and $CaTi_2O_4$ type $MgCr_2O_4$.

	Mg ₂ Cr	2 O 5		Cr ₂ O ₃			CaTi ₂	O ₄ type	
d(obs)	hkl	d(calc)	res(d)	hkl	d(calc)	res(d)	hkl	d(calc)	res(d)
5.213002	120	5.2092	0.0038						
4.818596	200	4.8161	0.00249						
4.2607							021	4.26542	-0.00472
3.800495	220	3.8018	-0.0013						
3.627271				012	3.63063	-0.00335			
3.391873							022	3.39019	0.00168
3.128075	230	3.13453	-0.00645						
3.098274	040	3.0965	0.00178						
2.945155	140	2.94791	-0.00276						
2.736519							110	2.73645	0.00007
2.664371				104	2.66113	0.00324	023	2.6687	-0.00432
2.634414							111	2.63318	0.00124
2.608906	240	2.6046	0.00431						
2.529234	330	2.53453	-0.0053						
2.498646	121	2.49312	0.00553						
2.479125				110	2.48263	-0.00351			
2.452912	201	2.44598	0.00694						
2.404155	150	2.39913	0.00503						

2 270002							040	0.07(0)	0.00205
2.378983							040	2.37603	0.00295
2.280135	221	2.27496	0.00517						
2.262742				006	2.25855	0.0042			
2.206014	250	2.20288	0.00314						
2.173841				113	2.17568	-0.00184			
2.157319							024	2.15576	0.00156
2.136896							042	2.13271	0.00419
2.117513							130	2.12186	-0.00434
2.100114	231	2.10439	-0.00428						
2.082865	041	2.09277	-0.00991				113	2.08663	-0.00377
2.070852							131	2.07261	-0.00176
2.013422	160	2.0185	-0.00507						
1.943569							32	1.94318	0.00039
1.910908							043	1.91299	-0.00208
1.893545	260	1.89738	-0.00383						
1.833987	151	1.83256	0.00142						
1.81373				024	1.81531	-0.00158	114	1.81238	0.00135
1.794802							025	1.79228	0.00252
1.771611							133	1.77265	-0.00104
1.759651	421	1.76074	-0.00109						
1.74406	530	1.74575	-0.00169						
1.700807							044	1.69509	0.00571
1.670563				116	1.67065	-0.00009			
1.649873	161	1.64516	0.00471						
1.615713	351	1.61375	0.00196	211	1.6137	0.00201	006	1.61267	0.00305
1.603345	600	1.60537	-0.00202				134	1.59515	0.0082
1.580593	511	1.58112	-0.00053	122	1.58043	0.00016	115	1.58002	0.00057
1.558837	620	1.554	0.00483				151	1.56189	-0.00305
1.541323	521	1.54384	-0.00251						
1.518497	550	1.52072	-0.00222						
1 503284							152	1 5042	-0.00092
1 486321	531	1 48716	-0.00083				102	1.0012	0.0007
1.472962	280	1.47396	-0.00099						
1 464203	200	1.11070	0.00077	214	1 46536	-0.00116			
1.707203				<i>4</i> 1 7	1.40550	0.00110			

1.429616	470	1.42588	0.00374	300	1.43335	-0.00373	135	1.42984	-0.00022
1.420615	002	1.41972	0.0009				153	1.42081	-0.00019
1.408371	560	1.40843	-0.00005						
1.391076	380	1.39458	-0.0035	125	1.39385	-0.00278	116	1.38935	0.00173
1.374701	461	1.37212	0.00258				202	1.37023	0.00378
1.363826	621	1.3632	0.00063				220	1.36823	-0.0044
1.334077							046	1.33435	-0.00027
1.317591							222	1.31659	0.001
1.302991	570	1.30315	-0.00016						
1.293531	232	1.29325	0.00028	1010	1.29247	0.00107			
1.285166				119	1.28742	-0.00226	136	1.28393	0.00124
1.270643	322	1.27081	-0.00017						
1.259425	740	1.25746	0.00197				223	1.25958	-0.00015
1.250375				217	1.2448	0.00558			
1.239382	0100	1.2386	0.00078	220	1.24132	-0.00193			
1.229906	191	1.22831	0.00159				204	1.23018	-0.00027
1.224463	402	1.22299	0.00147				155	1.22511	-0.00049
1.212235	721	1.21425	-0.00201	306	1.21021	0.00203			
1.202725	750	1.20291	-0.00018						
1.192875	252	1.19335	-0.00048	131	1.18803	0.00485			
1.166885	062	1.16978	-0.00289				028	1.17213	-0.00524



Appendix A5: Laser heating spot on the sample from both sides

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Appendix	A6:	Com	position	ot	different	samt	bles

Sample	MgO (%)	FeO (%)	Cr ₂ O ₃ (%)	Al ₂ O ₃ (%)	Others (%)
MgCr ₂ O ₄	20.9	-	79.1	-	-
FeCr ₂ O ₄	-	32.1	67.9	-	-
Natural Chromite	11	23	50	15	1

Ions	Radius (pm)
Mg ²⁺	86
Cr ³⁺	75.5
$\mathbf{Fe}^{2+}(\mathbf{ls})$	75
Fe ²⁺ (hs)	92
Al ³⁺	67.5
O ²⁻	140

Appendix A7: Ionic radius of different ions

Appendix A8: High pressure polymorphs of different spinel group minerals

Spinel	Polymorphs
MgCr ₂ O ₄	1. Cr_2O_3+MgO
	2. $Mg_2Cr_2O_5+Cr_2O_3$
	3. $CaTi_2O_4$ type structure
FeCr ₂ O ₄	1. $FeCr_2O_5+Cr_2O_3$
	2. $CaFe_2O_4$ -type structure
	3. $CaTi_2O_4$ -type structure
MgAl ₂ O ₄	1. $MgO+Al_2O_3$
	$2. Mg_2Al_2O_5 + Al_2O_3$
	3. $CaFe_2O_4$ type structure
	4. $CaTi_2O_4$ -type structure

CT-type MgCr ₂ O ₄					
Bond length (Å)				Bond angles (°)	
$Mg-O2^{\underline{i}} \times 2$	1.989(2)	$Cr-O3^{iii} \times 2$	1.958(1)	$O1^{\underline{v}i}$ –Cr1– $O3^{\underline{i}ii}$	172.4(1)
Mg–O 3^{ii} × 4	2.270(1)	Cr–O2	1.960(1)	02–Cr1–O3 ^{<u>iv</u>}	179.8(1)
MgO1 × 2	2.631(1)	$Cr-O3^{iv}$	2.016(2)	Cr1 ^{<u>vii</u>} –O1–Cr1 ^{<u>viii</u>}	89.58(4)
Average	2.290	$Cr-O1^{\underline{v}}\times 2$	2.023(1)	$Cr1^{\underline{vii}}$ - $O1^{\underline{vi}}$ - $C^{\underline{vi}}$	90.42(4)
<i>n_c</i>	4.93	Average	1.990	Cr1–O2–Cr1 ^{iv}	124.7(1)
BVS	1.90	n_c	5.95	$Cr1^{ix}$ –O3– $Cr1^{x}$	93.44(10)
		BVS	2.94	$Cr1^{ix}$ –O3– $Cr1^{iv}$	96.81(9)
mLd-type Mg ₂ Cr ₂ O ₅					
Bond length (Å)					
M1 site		M2 site		M3 site	
M1O1 × 4	2.010(2)	M2–O3 ^{\underline{x}} × 2	2.028(3)	M3–O2 ^{\underline{x}} × 2	2.012(2)
M1–O2 ^{\underline{x}} × 2	2.066(3)	M2-O4 × 4	2.117(2)	M3–O3 × 2	2.016(2)
Average	2.029	Average	2.088	M3–O4	2.062(3)
<i>n_c</i>	5.97	n_c	5.91	M3–O5 ^{<u>x</u>}	2.064(3)
BVS	2.43	BVS	2.07	Average	2.030
				n_c	5.97
				BVS	2.42
M4 site		M5 site			
M4O1 × 2	1.963(2)	M5–O1 × 2	2.099(2)		

Appendix A9 : Interatomic distances and angles in the structures of $CaTi_2O_4$ (CT)-type MgCr₂O₄ and modified ludwigite (mLd)-type Mg₂Cr₂O₅ (Ishii et al., 2015).
M4–O2 ^{<u>xi</u>}	2.011(3)	M5–O3 × 2	2.134(3)		
M4-O4 × 2	2.029(2)	M5-O5 × 2	2.190(3)		
M4-O5	2.035(3)	Average	2.141		
Average	2.005	n_c	5.93		
<i>n</i> _c	5.95	BVS	1.80		
BVS	2.59				
Bond angles (°)					
01–M1–O1 ^{<u>vii</u>}	90.6(1)	$M3^{\underline{ix}}$ - $O2^{\underline{vi}}$ - $M1^{\underline{xii}}$	121.2(2)	$M3^{ix}$ -O3-M3 ^{xv}	90.5 (1)
O1-M1-O2 ^x	97.32(7)	M1-O1-M1 ^{xii}	90.6(1)	M3–O5–M3 ^{<u>xii</u>}	90.2(2)
04–M2–O4 ^{<u>xii</u>}	84.85(8)	M4-O2-M1 ^{xiii}	95.85(3)	M4 ^{<u>vii</u>-O4-M3}	96.64(5)
03 ^{<u>x</u>} -M2-O4 ^{<u>xii</u>}	84.92(8)	$M1^{\underline{xii}}$ -O1-M4 $^{\underline{xiv}}$	96.15(5)	M3 ^{<u>xii</u>–O5–M4}	97.95(5)
O3 ^{<u>x</u>} -M3-O5 ^{<u>vii</u>}	169.29(9)	M2-O4-M2 ^{<u>vii</u>}	84.9(1)	$M4^{\underline{vii}}$ -O2-M3 ^{ix}	121.85(4)
O4–M3–O2 ^x	179.4(2)	M3-O4-M2 ^{<u>vii</u>}	92.0(1)	M4-O2-M4 ^{vii}	93.4(1)
02–M4–O4 ^{<u>xii</u>}	176.12(1)	M3 ^{<u>xv</u>} -O3-M2 ^{<u>ix</u>}	96.2(1)	M4-O4-M4 ^{vii}	89.5(1)
01 ^{<u>xi</u>} -M4-O5	172.0(2)	M4 ^{<u>vii</u>-O4-M2^{<u>vii</u>}}	92.16(3)		

Curriculum Vitae

Name:	Tauhid Belal Khan			
Post-secondary	[2004-2009] BSc. in Civil Engineering			
Education and	Bangladesh University of Engineering and Technology, Dhaka,			
Degrees:	Bangladesh			
	[2011-2013] MSc. in Advanced Clay Science			
	Université de Poitiers, Poitiers, France			
	[2013-2015] MSc. in Geophysics			
	University of Western Ontario, London, Ontario, Canada			
Honours and	[2013-2015] Western Graduate Research Scholarship			
Awards:	[2014] Canadian Exploration Geophysical Society travel bursary and book award			
	[2011-2013] Erasmus Mundus Scholarship			
Related Work	[2013-2015] Teaching Assistant and Research Assistant			
Experience	University of Western Ontario, London, Ontario, Canada			
	[2010-2011] Lecturer			
	University of Information Technology and Sciences, Dhaka, Bangladesh			

Publications:

Chemeda, Y.C., Christidis, G.E., **Khan, T.B.,** Koutsopoulou, E., Hatzistamou, V., Kelessidis, V.C., 2014. Rheological properties of palygorskite-bentonite and sepiolite-bentonite mixed clay suspensions, *Applied Clay Science, vol. 90, pp. 165–174*.

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