Geochemical and Petrological Characterizations of Peridotite and Related Rocks in Marquette County, Michigan

Andrew Lloyd Sasso

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GEOCHEMICAL AND PETROLOGICAL CHARACTERIZATIONS OF PERIDOTITE AND RELATED ROCKS IN MARQUETTE COUNTY, MICHIGAN

by
Andrew Lloyd Sasso

A thesis submitted to the Graduate College
in partial fulfillment of the requirements
for the degree of Master of Science
Geosciences
Western Michigan University
April 2016

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GEOCHEMICAL AND PETROLOGICAL CHARACTERIZATIONS OF PERIDOTITE AND RELATED ROCKS IN MARQUETTE COUNTY, MICHIGAN

Andrew Lloyd Sasso, M.S.
Western Michigan University, 2016

This study characterizes the following rock units in Marquette County, Michigan in terms of geochemistry and petrology: (1) Presque Isle Peridotite, (2) Deer Lake Peridotite, (3) Yellowdog Peridotite, and (4) Black Rock Point Gabbro. Analyses were conducted to determine if any petrological or geochemical relationships exist between these units, and to assess the potential of these units to host magmatic sulfide deposits.

The generated data and chosen geotectonic proxies indicate that Black Rock Point Gabbro and Deer Lake Peridotite crystallized from unrelated magmas, probably during the formation of the Great Lakes Tectonic Zone (2.7–1.85 Ga). No evidence was found which suggest that either unit hosts a magmatic sulfide deposit.

Investigation has also revealed a great deal of similarity between the Presque Isle and Yellowdog Peridotite units. Trace element comparisons, along with the use of geotectonic proxies (when considered in conjunction with Yellowdog Peridotite’s known age of 1107.2 Ma ± 5.7Ma) suggest the possibility that both units crystallized from a similar parent magma during the Midcontinent Rift event (1.1 Ga). Microprobe analysis revealed comparable Ni-depletion in both units. These findings suggest the possibility that Presque Isle Peridotite may be regarded as a prospective location for a magmatic sulfide deposit similar to that hosted by the Yellowdog Peridotite.
ACKNOWLEDGMENTS

I wish to thank my primary advisor Dr. Joyashish Thakurta for his patient guidance from the conception of this project, through its completion and for giving me the opportunity to peruse an advanced degree at Western Michigan University. I would also like to express my gratitude for the help and encouragement of the two other members of my thesis committee, Dr. Robb Gillespie and Dr. Mohamed Sultan. Additional, thanks are extended to the Western Michigan University, Graduate College for their financial support.

Further acknowledgement goes to Tony Boxleiter and Ben Hinks for their assistance in the field. And, to Tom Howe for supplying me with equipment.

Thanks also go out to Bob Mahin, of Lundin Mining, for allowing me to collect samples from the Eagle Mine, and for providing me with a geological map of the Eagle Mine area. Melanie Humphrey, of the DEQ for allowing me to access the Marquette core storage facility in order to collect samples. And, Dr. John Fournelle of University of Wisconsin, Madison, for his help with electron microprobe analysis.

To close, I wish to express my most sincere gratitude to my parents, Tony and Holly Sasso, for all of the support, and encouragement which they have provided throughout the undertaking of this project. I also wish to thank all of my friends who have lent their encouragement.

Andrew Lloyd Sasso
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<td>µm</td>
<td>Micrometer</td>
</tr>
<tr>
<td>BRP</td>
<td>Black Rock Point Area</td>
</tr>
<tr>
<td>DLP</td>
<td>Deer Lake Area</td>
</tr>
<tr>
<td>EA</td>
<td>Eagle East Intrusion</td>
</tr>
<tr>
<td>E-MORB</td>
<td>Enriched Mid-Ocean Ridge Basalt</td>
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<tr>
<td>EPMA</td>
<td>Electron Probe Micro-Analysis</td>
</tr>
<tr>
<td>EUG</td>
<td>Eagle Intrusion</td>
</tr>
<tr>
<td>ft</td>
<td>Feet</td>
</tr>
<tr>
<td>Ga</td>
<td>Giga annum (Billion years)</td>
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<tr>
<td>GLTZ</td>
<td>Great Lakes Tectonic Zone</td>
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<tr>
<td>GPS</td>
<td>Global Positioning System</td>
</tr>
<tr>
<td>ICP-MS</td>
<td>Inductively Coupled Plasma Mass Spectrometry</td>
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<tr>
<td>IUGS</td>
<td>International Union of Geological Sciences</td>
</tr>
<tr>
<td>km</td>
<td>Kilometer</td>
</tr>
<tr>
<td>LOI</td>
<td>Loss on Ignition</td>
</tr>
<tr>
<td>LREE</td>
<td>Light Rare Earth Element</td>
</tr>
<tr>
<td>m</td>
<td>Meter</td>
</tr>
<tr>
<td>Ma</td>
<td>Mega annum (Million years)</td>
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<tr>
<td>Meso-Prot.</td>
<td>Mesoproterozoic</td>
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<tr>
<td>Mg#</td>
<td>Magnesium Number</td>
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<tr>
<td>mi</td>
<td>Mile</td>
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<td>MORB</td>
<td>Mid-Ocean Ridge Basalt</td>
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<tr>
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<tr>
<td>OIB</td>
<td>Ocean Island Basalt</td>
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<tr>
<td>ppm</td>
<td>parts-per million</td>
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<tr>
<td>REE</td>
<td>Rare Earth Element</td>
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<tr>
<td>SEM</td>
<td>Scanning Electron Microscope</td>
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<td>Spn</td>
<td>Sphene</td>
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<tr>
<td>USGS</td>
<td>United States Geological Survey</td>
</tr>
<tr>
<td>wt. %</td>
<td>Weight percent</td>
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<tr>
<td>XRF</td>
<td>X-ray Fluorescence Spectrometry</td>
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LIST OF MINERAL ABBREVIATIONS

Ab  albite  (NaAlSi$_3$O$_8$)
Ac  acmite or aegirine (NaFeSi$_2$O$_6$)
An  anorthite  (CaAl$_2$Si$_2$O$_8$)
Ccp  chalcopyrite
Cpx  clinopyroxene
Fa  fayalite  (Fe$_2$SiO$_4$)
Fo  forsterite  (Mg$_2$SiO$_4$)
Fs  ferrosilite  (MgSiO$_3$)
En  enstatite  (MgSiO$_3$)
Hbl  hornblende  (Ca$_2$(Mg, Fe, Al)$_5$ (Al, Si)$_8$O$_{22}$(OH)$_2$)
Mag  magnetite
Ol  olivine  ((Mg, Fe)$_2$SiO$_4$)
Opx  orthopyroxene
Or  orthoclase feldspar  (KAlSi$_3$O$_8$)
Pn  pentlandite
Pl  plagioclase feldspar  (NaAlSi$_3$O$_8$ to CaAl$_2$Si$_2$O$_8$)
Px  pyroxene
Po  pyrrhotite
Wo  wollastonite  (CaSiO$_3$)
CHAPTER I
INTRODUCTION

Research Areas

Marquette County is located in the north central portion of Michigan’s Upper Peninsula (Figure 1). With an area of 1,821.5 square miles, Marquette is the largest county in the state of Michigan. This study focuses on four locations, and more specifically, four separate lithological units within this county. These four rock units have been mapped as peridotite by past researchers. They are located at: (1) Presque Isle, (2) Deer Lake, (3) Black Rock Point, and (4) the Eagle Mine site, on the Yellowdog Plains. These locations are shown in Figure 2.

Presque Isle

Presque Isle is a “tied” island in Lake Superior, connected to the mainland by a tombolo composed of lacustrine sediments. Located just north and east of the city of Marquette, Presque Isle has an exposed surface area of 323 acres and a maximum elevation of 724 ft near its center. Although originally designated as a government
A lighthouse reservation, Presque Isle has been designated as a park since being deeded to the city of Marquette in July of 1886. Today, Presque Isle is ringed by a paved road. It is Marquette County’s most popular destination for outdoor recreation and sightseeing.

Figure 2
Modified map of the four research areas and their surrounding geology. Adapted from Simms (1992).
**Deer Lake**

Deer Lake is a dammed lake located approximately 2 miles north of Ishpeming Michigan, and 12 miles west of downtown Marquette. The lake has a surface area of 950 acres, a surface elevation of 1,389ft, and a maximum depth of 35ft. There are a number of small islands scattered throughout the basin. Both low-lying, wetland areas and rugged, hilly terrain (dotted with outcrops) can been found in the lake’s immediate surroundings. Deer Lake is another of Marquette County’s more popular locations for outdoor recreational activities such as fishing, canoeing, and hiking.

**Black Rock Point**

Black rock point, also known as Black Rock Cape, is a small rocky point which juts due east into Big Bay (Lake Superior) off Salmon Trout Point. This point is located approximately 2.5 miles north of the unincorporated community of Big Bay, and approximately 26.5 miles north of Marquette. The point can be reached by traveling north from Marquette on county road 550 for 28 miles, then following county road Kf north for approximately 3 miles, followed by a hike of approximately 1 mile to the northeast. Due to its remote location and inaccessibility, the point is visited infrequently, and has never before been the subject of any geologic study.

**Yellowdog Peridotite-Eagle Mine**

The last location to be addressed by the project is the Yellowdog Peridotite located at the Eagle Mine site near the center of the Yellowdog Plains. This area is located approximately 9 miles southwest of Big Bay, and approximately 27 miles
northwest of Marquette. It consists of a solitary outcrop, known as Eagle Rock, which is now surrounded by the surface infrastructure of the Eagle Mine operation.

**Peridotite**

Peridotite is a plutonic, ultramafic rock type characterized by a mineralogical composition dominated by 40-100 vol% Olivine (Wooley, 1996). Other mafic minerals such as pyroxenes and amphiboles account for most of the rock’s remaining volume. Additionally, minerals such as plagioclase, garnet, chromite, and spinel often comprise a minor fraction of the rock volume. These rocks are typically coarse grained and dark colored. Peridotites are believed to be the dominant rock type found in Earth’s upper mantle; however, they are relatively rare at the surface. The extrusive equivalent of peridotite is komatiite.

Peridotite is classified based on its mineralogical composition, and constitutes all rocks composed

**Figure 3**
IUGS classification tables for ultramafic rocks. Peridotite shown in yellow. (Woolley, 1996)
of a minimum of 40 vol% olivine. Dunite is a peridotite composed of a minimum of 90% olivine, with minerals such as pyroxene, chromite and spinel constituting its remaining bulk. Harzburgite is a peridotite composed of 40-90 vol% olivine, whose pyroxene fraction is at least 95 vol% orthopyroxene minerals. Similarly, wehrlite is composed of 40-90 vol% olivine, with a pyroxene fraction represented by no less than 95 vol% clinopyroxene minerals. Lherzolite is the most common variety of peridotite, it is composed of 40-90 vol% olivine, and a pyroxene fraction ranging between 95 vol% clinopyroxene, and 95 vol% orthopyroxene. Ultramafic rocks composed of less than 40 vol% olivine include pyroxenite and hornblendite. These ultramafic classes have compositions dominated by pyroxene and amphibole. Pyroxenite and hornblendite are subdivided, in the same fashion as peridotites, based on the chemistry and abundance of their dominant mafic minerals. Figure 3 is the standard classification table for ultramafic rocks developed by the International Union of Geological Sciences, it provides a visual description of the ultramafic classification scheme.

All peridotites found at or near The Earth’s surface originate from one of two sources: (1) Ophiolites or (2) Cumulates.

**Ophiolites**

Peridotites may have formed within The Earth’s upper mantle and subsequently been emplaced within the upper crust by tectonic processes (such as accretion during subduction). Such sequences are termed ophiolites. Ophiolites are defined as thrust sheets of ancient oceanic crust and upper mantle rock which has been abducted over, and accreted onto the edge of the continental crust during the course of orogeny (Dalik et al., 2003). Ophiolites are easily identified by their standard lithological sequence, known as a
Typical Ophiolite Suite. This sequence, from the top down is: (1) Deep sea sediments, (2) pillow basalt, (3) sheeted dike complex, (4) isotropic gabbro, (5) layered gabbro, (6) Mantle peridotite (Coleman, 1977).

One example of this sequence is shown in Figure 4.

Peridotites found in ophiolites formed as part of the upper mantle but have since been tectonically emplaced within the upper crust. Typical peridotite of this type can be found at Bay of Islands, Newfoundland (Figure 4) and at the Semail Ophiolite, Oman.

**Cumulates**

Peridotites may also be formed as cumulates. Cumulates are the result of differential crystallization and subsequent accumulation of olivine and pyroxene from a mafic or ultramafic melt which formed deep within the Earth and has intruded the crust. The deep origin of such a mafic melt generally results from convective melting due to a mantle plume (Wagner et al., 1960). When these magmas intrude into the crust, pressure
and temperature decrease, and this causes crystallization to begin. High temperature minerals such as olivine and pyroxene will be the first to crystallize. They will settle due to their high density and accumulate at the base of the magma chamber. The exact temperature at which crystallization occurs depends on the pressure, and the FeO/MgO ratio of the parent melt. Continued accumulation of mainly mafic minerals will result in the formation of a substantial ultramafic pluton. Remaining parent melt, having been depleted of mafic material, will eventually cool and crystallize, resulting in a zone of the same pluton less mafic than its associated cumulate. Figure 5 depicts this process.

Typical peridotites of this type can be found in the Bushveld Igneous Complex, South Africa and the Bovine Igneous Complex, Baraga County, Michigan.

![Figure 5](image)

**Figure 5**
Graphical representation of cumulate rock formation.

Olivine and pyroxene, as discussed above, are the chief mineral components of peridotite and crystallize under conditions of high temperature and pressure. Once
emplaced within the crust (even more so when exposed at the surface) these high pressure minerals quickly become unstable. Interaction with hydrothermal fluids near the surface leads to hydration of these primary ferromagnesian minerals. This process results in their alteration to secondary serpentine group minerals which are stable at, and near, atmospheric temperature and pressure. This low temperature alteration process (0-500°C), is known as serpentinization. Other common alteration minerals include brucite, chlorite, talc, magnesite and calcite (Evans, 1977).

Magmatic Sulfide Deposits

Magmatic sulfide deposits can be defined as “substantial” concentrations of sulfide minerals in ultramafic or mafic rocks, which have been derived from immiscible sulfide liquids (Foose et al, 1986). These deposits are important sources of copper, nickel and platinum group elements. They are mined worldwide.

Magmatic sulfide deposits can be divided into two basic categories: (1) Those rich in sulfide containing high concentrations of nickel and copper, and (2) those relatively sulfide poor, but containing high concentrations of platinum group elements (Naldrett, 1999). The Eagle Ni-Cu-PGE deposit, located in Marquette County, is of the former variety. This will be the focus of this study.

Formation of economic magmatic sulfide deposits is dependent upon three key factors (Naldrett, 1999):

(1) The host magma must become saturated in sulfide, allowing for the segregation of an immiscible sulfide liquid. Because the segregation of a sulfide liquid and its subsequent
basal accumulation is not normally observed during the cooling and crystallization of mafic magmas, sulfide saturation of the parent melt must be required for such an occurrence. Mafic magmas are generally not sulfide saturated when they form in the upper mantle, or begin crustal intrusion; therefore, a significant source of sulfur must be encountered, and allowed to contaminate such a melt, for sulfide saturation to be reached.

(2) Sulfides must be allowed to react with a volume of magma sufficient for the concentration of chalcophile elements such as Fe, Ni, Cu, and platinum group elements, to an economic level. Economic magmatic sulfide deposits will only form when the parent melt is of sufficient volume and has a metal-rich chemistry. The parent melt must also continue interaction with the immiscible sulfide liquid even after segregation in order for the sulfide liquid to attain economic levels of chalcophile element enrichment. Ultramafic melts, which contain the highest fraction of metallic chalcophile elements, and lowest silica concentration, best meet these conditions, and are therefore the ideal parent magma for magmatic sulfide genesis. Mafic melts are likewise considered excellent hosts for magmatic sulfide formation. This explains why magmatic sulfide deposits are invariably found in association with ultramafic, and mafic rock units.

(3) Once formed, crystalline sulfides must become concentrated in significant abundance, in order to constitute an ore body.

Schulz et al. (2010), states that magmatic sulfide deposits are generally related to a range of olivine-bearing, mantle-derived magmas, which may result in a variety of lithologies. Parental magmas of magmatic sulfide hosts, are typically under saturated in sulfur and relatively enriched in Cu, Ni, and PGEs. These parent melts are also enriched in other strongly incompatible elements such as K, P, Ba, Sr, Pb, Th, Nb, and light REEs.
Marquette County: Geologic Background

Great Lakes Tectonic Zone

The Great Lakes Tectonic Zone (GLTZ) is a 1,400 km long crustal boundary which extends from present-day South Dakota, eastward to the vicinity of Sudbury Ontario. It passes directly through Marquette County, Michigan, where it is well exposed. The GLTZ separates the “Northern Complex” greenstone, granite terrain of the southern Superior Craton, to the north side, from “Southern Complex” gneiss terrain to the south. It is interpreted as a Late-Archean continent-continent collision zone formed during the Kenoran Orogeny (2.7-2.5 Ga). The Minnesota River Valley terrane, during this time, collided with, and overrode, the Superior Craton. This resulted in the formation of the GLTZ, and also produced widespread deformation and metamorphism in the vicinity of the boundary. The initial formation of the GLTZ was followed by a period of extension (2.4-2.1 Ga). Following this series of events, a second major episode of compression occurred during the Penokean Orogeny (1.85-1.83 Ga), which occurred to the southwest of the main event. Even today, minor activity is infrequently observed along the GLTZ. Documented earthquakes have occurred as recently as 1975. The GLTZ in Marquette County, appears as a 2 km wide zone of northwest-striking, metamorphic rock, separating the distinctly different lithologies of the northern and southern complexes (Sims, 1991). Figure 6 shows a simplified map of the GLTZ as it extends through Minnesota, Wisconsin, and western Upper Michigan (including Marquette County).
Figure 6
Simplified map of Great Lakes Tectonic Zone.
From Sims et al. (1993).
The Midcontinent Rift

The Midcontinent rift is a failed continental rift system which formed 1.1 billion years ago during the Mesoproterozoic. It is conventionally thought that formation of the rift was caused by a mantle plume. Creating a triple junction with its nexus at Lake Superior (Stein et al., 2014). Only two arms of the rift are well developed. One arm extends southwest from Lake Superior, and can be followed to its point of termination in present-day Kansas. The second arm extends southeast where it is covered by sediments of the Michigan Basin. The poorly developed third arm extends northward where igneous materials associated with its formation can be observed in the vicinity of Lake Nipigon in present-day Ontario, Canada. Although the cause of the rift’s failure is not known, compression to the southeast due to the Grenville Orogeny is one possible explanation (Schmus, 1985). Figure 7 shows a generalized diagram of the rift system.

Rocks in the Lake Superior region associated with the rift comprise a series of plutonic, volcanic, and clastic sedimentary units dating from 1.0 Ga to 1.2 Ga. These are collectively referred to as the “Keweenawan Supergroup”. The primary feature in Marquette county associated with the rift is a massive swarm of mafic dikes which trend east-west, parallel to the rift axis. These features formed by intrusion of molten material.
into the crust along the margins of the rift. Figure 8 shows a generalized interpretation of the relationship of these features to the rift system.

**Purpose**

Previous researchers have mapped peridotite rock units at four locations across Marquette County, Michigan. These units are found at: (1) Presque Isle, (2) Deer Lake, (3) Eagle Mine, and (4) Black Rock point. The locations of these sites are shown in Figure 2.

Exploration by Kennecott in 2005 discovered a world class magmatic sulfide deposit hosted within the Yellowdog Peridotite. This discovery has since been developed into the current Eagle Mine site. Discovery of a magmatic sulfide deposit in association with one of the Marquette County peridotite units has raised two significant questions.
Could an as yet undiscovered magmatic sulfide deposit, similar to that found in association with the Yellowdog Peridotite, exist in association with any other Marquette County peridotite unit? Secondly, can a petrological or geochemical relationship be established between any of Marquette County’s peridotite units?

Both the Yellowdog Peridotite and associated magmatic sulfide deposit have been studied extensively since the 2005 discovery. Radiometric dating of the peridotite returned an age of 1107.2 ± 5.7 Ma (Ding et al., 2010). These results imply that the peridotite, along with its associated magmatic sulfide deposit, formed during the midcontinent rift event.

Although a number of previous studies have been conducted out at both the Deer Lake, and the Presque Isle peridotite units, no radiometric dates have been obtained for either unit. Previous researchers have collected and analyzed an abundance of petrological and geochemical data for all but the Black Rock Point unit. Despite this effort, no formal petrological or geochemical comparisons have been established to correlate the units.

These two questions (above) are significant, not only due to their economic implications, but also due to advancing the collective understanding of geologic history in Marquette County. It has been the goal of this study to answer these questions through the characterization, and subsequent comparison, of Marquette County’s peridotite units in terms of both petrology and geochemistry. A digital geologic map of each peridotite has also been created. This was done to fully visualize the scope and surrounding geology of each peridotite unit.
Previous Work

Marquette County is perhaps best known for the Marquette Iron range. Since its discovery in 1844, by William A. Burt, the range has been continuously mined. Due to the presence, and the substantial value of this large iron range, the area has been the focus of extensive mineral exploration during the past 170 years. This exploration has led to the discovery of numerous economic minerals and native element deposits. More than 380 mines and prospects have been established in Marquette County.

The most recent mine established in Marquette County is the Eagle Mine, located on the Yellowdog Plains. This mine is owned and operated by Lundin Mining Corporation. The mine’s primary target resources are nickel and copper in the form of magmatic sulfides. Lundin also expects to recover minor quantities of platinum, palladium, and cobalt. The primary ore deposit is estimated to be approximately 4.1 million metric tons. It is hosted by the Yellowdog Peridotite, an ultramafic intrusion dating to 1.1Ga, the time of the Midcontinent Rift event. Ore being extracted at the mine has been found to contain 3.6 wt. % nickel, and 2.9 wt. % copper. Nearly 8 billion U.S. dollars’ worth of ore is expected to be extracted during the project’s projected 8 year mining life (Owen & Meyer, 2014). The Eagle mine began production on September 23, 2014. Eagle’s discovery and development has set a precedent for economic mineral deposits associated with peridotite intrusions in Marquette County.

Klanser et al. (1979), characterized this rock unit, prior to the discovery of economic ore deposit associated with the Yellowdog Peridotite. Since then, numerous other papers and technical reports have been published regarding the geology of the area,
primarily by Kennecott Exploration, Rio-Tinto, Lundin Mining Corporation, and the consulting firm Wardell-Armstrong.

Previous geologic research has also taken place at both the Presque Isle and Deer Lake sites. The earliest identification of peridotite at Deer Lake was made by Van Heise in 1895. Rossell et al. (1983) also conducted a detailed study of the Deer Lake Peridotite which was first to identify two petrologically distinct rock types within the unit.

Presque Isle has been the subject of numerous theses and research projects since the late 19th century. The earliest attempt to characterize Presque Isle’s geology was made by Wadsworth (1884). Wadsworth’s work formed the foundation for all subsequent work on the island’s geology. There is a section on the Presque Isle Peridotite that appears in the 1895 paper “Preliminary Report on the Marquette Iron-Bearing District of Michigan” by Van Heise. The most recent detailed report on Presque Isle’s geology was written by Lewan et al. (1972). A less detailed follow up to Lewan’s work was conducted by Rick Lantz (1982).

No previous research has been conducted in the vicinity of Black Rock Point, and very little mention has been made of the site in any previous publications or unpublished works. The point was mapped as serpentinitized peridotite by Case and Gair, 1985; however Black Rock Point is never specifically mentioned in the report. During mapping for this study, it was quickly discovered that no peridotite actually exists at Black Rock Point. The area is actually composed of two main rock types separated by diabase dikes of Keweenawan age. The southernmost and most prominent of the major types, is a large mafic body of gabbro. The other main rock type present is a felsic, heavily veined gneiss.
Presque Isle Peridotite

Serpentinized peridotite at Presque Isle, makes up approximately two thirds of the island’s 323 acre area. This peridotite shows varying degrees of alteration, but appears to be moderately to heavily serpentinized at most locations. It is likely that this peridotite unit dates to the late Archean or early Proterozoic; however, its composition and variable alteration makes determination of its exact age challenging. Gair and Thaden (1968) have hypothesized that the peridotite could have formed as late as 1.1 Ga in the early stages of the Midcontinent Rift event.
Jacobsville Sandstone unconformably overlies the peridotite and constitutes the remaining third of Presque Isle. The Jacobsville Sandstone is notable being the youngest unit in the Keweenawan series.

Lewan et al. (1972), noted the peridotites at this site form a tear drop shaped mass approximately 1500 meters long and 700 meters wide. Peridotite forms the major basement rock at this site. It is heavily fractured and often cut by veins of serpentine, or carbonate minerals emplaced by hydrothermal activity. Sulfides such as chalcopyrite and galena are visible in these veins at some locations. Prominent peridotite outcrops exist along Presque Isle’s northern and eastern shoreline, although the majority of the peridotite is covered by either glacial material or Jacobsville Sandstone. These outcrops, in some areas, form cliffs greater than 100 feet in height. Jacobsville Sandstone is a common unit across Marquette County’s Lake Superior shoreline; however, near its contact with the peridotite at Presque Isle, the unit displays an atypical appearance. The Jacobsville Sandstone and its basal conglomerate in particular, overlie the peridotite in a nonconformity. They are heavily cut by carbonate veins similar to those found in the peridotites. The basal conglomerate consists of angular fragments of chert, cemented in a matrix of fine quartz sand. It is likely that the conglomerate is derived from the underlying alteration zone (Lewan, 1972). Locally, the sandstone is seen to contain rounded cobbles of the peridotite (Lantz, 1982).

Peridotite is not seen in contact with any major rock units other than the Jacobsville as its exposure is surrounded by Lake Superior. However it is assumed by observing the geology of the surrounding islands and mainland, that the unit is hosted by the Compeau Creek Gneiss or basement granite on the “Northern Complex”.
Deer Lake Peridotite

The Deer Lake Peridotite is located within the Ishpeming Greenstone Belt. It is bounded to the south by the east west trending Great Lakes Tectonic Zone. The unit is approximately 8km long and 1km wide. Deer Lake Peridotite displays variable degrees of alteration, but is generally heavily to entirely serpentinized. It is hosted by the Kitchi Schist which consists of: (1) A lower mafic member of metamorphosed basalts, diabases, and gabbros, and (2) upper member of andesites, coarse dacitic pyroclastics, berrecias,
and fine grained tuffs. The Kitchi Schist is overlain by the Mona Schist which has been dated to 2.7 Ga. This unit consists of pillow basalts and felsic pyroclastic rocks. There has been some debate as to whether the sequence between the Kitchi and the Mona is conformable. (Morgan and DeCristoforo, 1974 Rossell, 1983). The Deer Lake Peridotite is truncated by the Great Lakes Tectonic Zone on its south western margin. This can be observed in Figure 2. Truncation of the unit by the Carp River Falls Shear Zone, which represents the northern margin of the GLTZ in Marquette County, suggests that it must have been emplaced prior to, or during, formation of the GLTZ (2.7-1.85 Ga).

**Black Rock Point Gabbro**

![Geological map-Black Rock Point showing sample locations, samples collected during May 2014 and May 2015.](image)

*Figure 11*
Geological map-Black Rock Point showing sample locations, samples collected during May 2014 and May 2015.
This small, remote point which juts into Lake Superior just north of Big Bay is composed of two main rock units, one mafic, and one felsic as described earlier. The felsic unit has been cut by two mafic dikes which are probably of Keweenawan age. Other than the relatively small exposure of Black Rock Point, the entire unit is overlain by a thick layer of Jacobsville Sandstone which forms cliffs greater than 50 ft high. Field mapping during May 2015 led to the discovery of additional outcrops of gabbro and granite located above the sandstone cliffs.

Yellowdog Peridotite

The Yellowdog Peridotite is located near the center of the Yellowdog Plains, an area of Pleistocene glacial outwash. This ultramafic intrusion is associated with the Mesoproterozoic dike swarm found trending east-to-west across Marquette County and the surrounding region. The Yellowdog Peridotite formed during the Midcontinent Rift event (1.1 Ga). This is confirmed through the use of U-Pb baddeleyite age dating (Ding et al., 2010) which determined an age of 1107.2 Ma ±5.7 Ma. This peridotite occurs as distinct western and eastern intrusions, referred to as “Eagle” and “Eagle East”. Although the two intrusions are conventionally thought to be connected beneath the surface, exploration drilling has thus far failed to confirm this. The Eagle East intrusion rises the surrounding glacial outwash plain to form a prominent knob. Approximately 650 m to the west of this outcrop is the western or “Eagle” intrusion. This portion of the peridotite intrusion has no surficial outcrop. Both intrusions have a strike length of approximately 530 m and dip steeply to the north. These intrusions are keel shaped and range from approximately 75m wide near the surface to 5 m in width at a depth of 300 m. Both
intrusions host world class magmatic sulfide deposits rich in nickel and copper, along with notable amounts of platinum group elements (PGE). The deposit is actively mined as Lundin Mining Corporation’s Eagle Mine (Owen and Meyer, 2013).

The Yellowdog Peridotite is situated at the east most end of the Baraga Basin, the northern most basin of Paleoproterozoic sediments in Michigan (Owen and Meyer, 2013). The host (basement) rock is a gneiss-greenstone complex dating to more than 2.5 Ga. It is overlain by the Marquette Range Supergroup which dates to approximately 2 Ga (Klasner et al., 1979). These rock units are shown in map form in Figure 12.
Figure 12
Map showing locations of the Yellowdog Peridotite and its surrounding rock units. Adapted form (Dunlop, 2013) by Bob Mahin of Lundin Mining.
CHAPTER III
METHODS

Field Methods

Field work was conducted during May, 2014 (with the exception of the Eagle site). Various hand samples of both the targeted rock units, and the surrounding lithologies, were collected from each area. Coordinates for each sample location were recorded using a handheld GPS device. Tables 1-4 list samples collected from each field area by sample number, rock type, and sampling location. MQT samples are from Presque Isle, DLP samples are from Deer Lake, BRP samples are from Black Rock Point, EA samples are from the Eagle East intrusion, and EUG samples are from the Eagle intrusion. Local field mapping was also conducted to better define their outcrops and delineate boundaries between major rock units. Important geologic contacts and other notable features such as veins, dikes and areas of sulfide mineral occurrence was also mapped and recorded.

Additional field work was conducted at all research areas during May, 2015. Mapping continued at the Deer Lake site to better delineate the extent of the Deer Lake Peridotite. Samples were collected from beneath the eastern intrusion of the Yellowdog Peridotite from within the Eagle Mine. Additional hand sample collection and minor map adjustments also occurred at the Presque Isle, and Black Rock Point sites.

Table 1: Yellowdog Samples

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<th>Sample Number</th>
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<td>87° 52’ 57.60&quot;W</td>
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<td>Core</td>
<td></td>
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<td>Core</td>
<td></td>
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<td>Rock Type</td>
<td>Latitude</td>
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Table 3: Deer Lake Samples

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Table 4: Black Rock Point Samples

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Petrology and Petrography

After the 2014 field season, ten hand samples were selected, cut and sent to Spectrum Petrographics Inc. for thin section preparation. Samples were selected for each site to be representative of the major rock units found at those locations. Five of these
samples represented several different rock units from the various sample locations around Presque Isle, two were from distinctly different peridotite samples collected in the vicinity of Deer Lake, and three were from the Black Rock Point site (two samples were gabbro and one was a mafic dike).

This process was repeated for samples collected during the 2015 field season. Thirteen samples were selected, cut and sent to Vancouver Petrographics Ltd. for thin section preparation.

All thin sections collected from both the 2014 and 2015 seasons were analyzed in plane polarized, cross polarized, and reflected light using a Leica DM 750P petrographic microscope. This petrological analysis was conducted to create a detailed description of each unit’s mineralogical and textural characteristics. A list of this sections is available in Table 5. MQT=Presque Isle, BRP=Black Rock Point, EA=Yellowdog (Eagle Mine), DLP=Deer Lake.

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<tr>
<td>MQT-14-003</td>
<td>peridotite</td>
</tr>
<tr>
<td>MQT-14-007</td>
<td>highly altered peridotite</td>
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<tr>
<td>MQT-14-011</td>
<td>peridotite</td>
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<td>MQT-14-014</td>
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<tr>
<td>MQT-14-016</td>
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### Table 5: Continued

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<tr>
<td>DLP-1784ft</td>
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**Geochemistry**

Samples from each study area were also selected to undergo geochemical analysis. Geochemical data reported in this study are the result of three separate analytical techniques employed by two separate laboratories.

Eleven Samples were sent to Geoscience Laboratories, located in Sudbury Ontario. Samples MQT-15-002, BRP-14-005, EA-15-001, and DLP-1784 were analyzed using X-Ray Fluorescence Spectrometry (XRF). These samples were first crushed and pulverized to a grain size of <75µm, before being mixed with a flux, and heated to approximately 1000°C. This resulted in the production of a fused glass disc. This disc, which is representative of the sample’s original chemistry, is free of any mineral structure, and is ideal for XRF analysis.
In preparation for analysis employing Inductively Coupled Plasma Mass Spectrometry (ICP-MS) samples, MQT-15-002, MQT-14-003, MQT-14-011, 14EA331H1141.9m, 04EA054 86.05m, EA-15-001, EA-15-003, BRP-15-012, BRP-14-005, DLP-15-006, and DLP-1784 were first crushed and pulverized before being fully dissolved in a closed vessel multi acid digestion. The resulting solution was then ready to be analyzed using ICP-MS.

Electron microprobe analysis (EPMA) was conducted at the University of Wisconsin-Madison’s Eugene Cameron Electron Microprobe Lab, under the supervision of Dr. John Fournelle. The instrument used was a Cameca SX50 microprobe. Polished thin sections BRP-14-005, MQT-14-003, EA-15-001, and EAUG0012-86.43m were first sterilized and carbon coated to increase their conductivity. Next, the slides were examined using a scanning electron microscope (SEM). This device allowed for observation of the thin sections in much greater detail and at higher magnification than is possible with a standard petrographic microscope. Following (SEM) imaging, the slides were loaded into the electron microprobe. The Electron microprobe was able to target the most pristine grains of the primary minerals remaining in each sample. The Deer Lake Peridotite was excluded from this portion of the study because a thin section containing a sufficient fraction of primary minerals could not be produced.

Geological Mapping

Data collected during field work were first compiled and digitized along with data collected from previous work at each site. These digitized data include unit boundaries, important structural features, outcrop locations, hand sample locations, and unit
descriptions. This includes information for both the units directly addressed in this study and those in their immediate vicinity.

Digitization was accomplished through the use of ESRI-Arc GIS software. Digitized data were then incorporated into the appropriate maps within the GIS software. Layers containing the relevant data could then be used as overlays atop the topographic base map for each study area. Then, by applying a digital elevation model available from the USGS online database, a new geological map could be created for each research site. The resulting maps contain all data collected during the field portion of this study. It also includes all relevant data recorded by previous researchers, making these maps an ideal accompaniment to the petrological and geochemical analyses and interpretations which follow.
CHAPTER IV
RESULTS

Petrographic Description

Presque Isle Peridotite

Presque Isle Peridotite is best described as a medium-to fine-grained lherzolite (peridotite composed of ≥ 40 vol% olivine, and whose pyroxene component is made up of both orthopyroxene and clinopyroxene). Since its formation, the unit has undergone significant alteration. Remaining primary minerals range from <5% to >30% depending upon sampling location. The unit’s altered fraction is generally composed of serpentine in the form of rounded pseudomorphs after olivine (these are generally recognizable as rounded crystals displaying irregular cracks), and pyroxene-bastite (serpentine pseudomorphs after clinopyroxene). Other secondary minerals such as hematite, magnetite, goethite, carbonates, and quartz constitute 1-2% of the unit. Sulfides such as pyrite and galena also account for 1-2%.

Presque Isle Peridotite, in outcrop, weathers to a brownish green color. Hand samples from Presque Isle Peridotite display a greenish black coloration on fresh surfaces, and a phaneritic texture. However, individual crystals are difficult to distinguish in areas where the rock displays higher degrees of alteration. This is due to the presence of numerous hydrothermal veins filled by serpentine, quartz, calcite, carbonate minerals, or sulfide minerals (such as pyrite and galena).
Figure 13
Photo of sample MQT-15-002
- Hand Sample-Lightly serpentinized Presque Isle Peridotite.
- Note visible phaneritic texture and brownish weathering rind.

Figure 14
Field photo from eastern side of Presque Isle, showing Presque Isle Peridotite as it appears in outcrop.
Thin sections MQT-14-001, MQT-14-003, MQT-14-011, and MQT-15-002 are Presque Isle Peridotite, and display approximately 60-98% alteration minerals. Alteration is primarily observed in the form of serpentine after olivine pseudomorphs with clinopyroxene-bastite filling the interstitial areas is the unit’s principal mineral assemblage. Where primary minerals that are still present are composed primary of olivine surrounded by pyroxene grains. The unit exhibits a cumulate texture consisting of subhedral to euhedral olivine with interstitial pyroxene. There is a poikilitic texture that can be observed in some cases, with olivine chadacrysts are surrounded by pyroxene oikocrysts. This texture is shown in Figure 15. Primary minerals can best be observed in thin sections MQT-14-003, and MQT-14-011. Small crystals of spinel and sulfide minerals can also be observed in thin sections of this rock type regardless of alteration level. Sulfide Minerals are commonly observed as inclusions within olivine crystals as shown in Figures 17 and 18.

Thin section MQT-15-002 shows significant sulfide mineralization in the form of pentlandite, and chalcopyrite, which has crystallized in association with magnetite, as seen in Figure 18. Pentlandite is distinguished by its light yellowish cream reflection color, from its supergene equivalent, violarite, which reflects grey to pink (Pracejus, 2008).
Figure 15
Photomicrograph of MQT-14-003
Olivine chadacrysts surrounded by pyroxene oikocrysts.

Figure 16
Photomicrograph of MQT-14-001
Serpentine after olivine pseudomorphs.
Figure 17
Reflected light photomicrograph of MQT-14-011. Sulfide inclusions in olivine indicated by red arrows.

Figure 18
Deer Lake Peridotite

Deer Lake Peridotite, in its pre alteration state, would have been described as a medium grained peridotite. However, the majority of the unit has been highly altered. Collecting samples, at some locations, which retain their original crystallinity and texture is possible. However, the unit’s primary minerals have invariably been replaced by serpentine pseudomorphs of olivine and pyroxene. Various other secondary and alteration minerals are also present in smaller amounts. Based on observations of these secondary mineral structures, it is apparent that the unit’s original mineral assemblage was once dominated by olivine and pyroxene.

Figure 19
Photo of sample DLP-15-006.
Hand sample of low alteration Deer Lake Peridotite.
Deer Lake Peridotite, in hand sample, appears to be fine to medium grained and is greenish to black in coloration. The unit typically weathers to a brown-grey color where it is exposed at the surface. Some samples are highly altered, making their crystallinity difficult to distinguish.

Thin sections, DLP-15-006, DLP, 1784ft, and DLP-562ft are Deer Lake Peridotite. Rossell (1983), notes that peridotite samples collected at Deer Lake appear to be of two varieties, both of which display nearly complete serpentinization. Type 1, is commonly found within outcrops of the peridotite. Samples of this rock type show no foliation; rather, they retain a relic igneous texture (observed in thin section, DLP-15-006). This relic igneous texture is characterized by pseudomorphs of serpentine after olivine, identified by rounded crystals and irregularly cracked texture. Serpentine after pyroxene pseudomorphs (known as bastite) are also present. Other minerals such as
magnetite, various carbonates, and talc comprise a smaller fraction of the unit. Similar to the Presque Isle Peridotite, serpentine after olivine pseudomorphs are often enclosed in anhedral bastite. Observation of these relic textures permits the interpretation, that in its pre-alteration state, the unit would have displayed a cumulate texture. There is a relic poikilitic texture observed in some areas, here pyroxene oikocrysts enclosed olivine chadacrysts. These pseudomorphs and associated textures, are exhibited in Figures 21 and 22. Small grains of spinel and sulfide minerals, can also be observed throughout thin sections of the Deer Lake Peridotite however, they account for only 1-2% of the rock unit.

Figure 21
The second variety, or Type 2, of peridotite observed at Deer Lake has been observed only in drill core samples. These were originally obtained by Callahan Mining Company in the vicinity of the Ropes Gold Mine, and collected from the Michigan Department of Environmental Quality’s core repository located in Harvey, Michigan. Type 2 are highly foliated. Serpentine is the most abundant mineral. Other minerals such as magnetite, various carbonates, and talc comprise a smaller fraction of the unit. The high degree of foliation has drastically distorts, or completely destroys any relic igneous texture. This makes it impossible to determine (through petrographic means) whether the primary mineralogy and texture of Type 2 samples would could been similar to that interpreted for Type 1 samples.

Figure 22
Photomicrograph of DLP-15-006 Bastite oikocrysts (Px) surround serpentine after olivine chadacrysts (Ol).
Black Rock Point Gabbro

Petrographic analysis of the Black Rock Point Gabbro allows for its classification as a medium-to-coarse grained gabbro. The unit is composed primarily of plagioclase feldspar and clinopyroxene. Sphene is also found in small quantities. This gabbro shows a substantial degree of alteration resulting in amounts of chlorite, quartz, and amphibole being present.

In hand sample, Black Rock Point Gabbro is bluish green in color and is medium to coarse grained in texture. It weathers to an unusual reddish tone where the unit is exposed to the surface.

Figure 23
Black Rock Point Gabbro.
Photo of hand sample BRP-14-005.
Thin sections BRP-14-005, BRP-14-006, and BRP-15-012 are Black Rock Point Gabbro. Petrographic analysis reveals the units to primarily have a hypidiomorphic texture in which the majority of crystal phases are subhedral to anhedral. Plagioclase crystals are subhedral, easily recognized by their: (1) First order interference color, (2) lath shape, and (3) albite twinning. Pyroxene can be identified by its: (1) High refractive index, (2) second order interference color, and (3) inclined extinction. Sphene sparsely occurs throughout the unit as anhedral crystals of varying size. It can be identified by its very high positive relief, and cleavage. Small plagioclase laths are commonly found either partially or fully enclosed by clinopyroxene, giving the unit a subophitic to ophitic texture.

Figure 24
Black Rock Point Gabbro outcrop.
Figure 25
Photomicrograph from BRP-14-005 showing plagioclase and clinopyroxene crystals. Plagioclase lath indicated by red arrow is partially enclosed within clinopyroxene.

Figure 26
Photomicrograph form BRP-14-005. Sphene (Spn) is visible at center of image.
Yellowdog Peridotite

Yellowdog Peridotite found in both the eastern and western intrusions, is a medium to coarse-grained feldspathic lherzolite. This unit is composed primarily of olivine with clinopyroxene, orthopyroxene, and plagioclase feldspar. Yellowdog Peridotite is partially serpentinized, and alteration of both olivines and pyroxenes can be observed. The unit, a medium-to coarse-grained plagioclase lherzolite, has a composition of ≥40% olivine, contains a significant fraction of both orthorhombic and monoclinic pyroxene. Substantial quantities of plagioclase feldspar can also be observed.

Yellowdog Peridotite, In hand sample, is dark grey to greenish colored and has a mottled white and black appearance due to the presence of plagioclase. The rock, in outcrop, is weathered brown to red due to of pyrrhotite (an iron sulfide) hosted within the unit.

Figure 27
Photo of sample EA-15-002.
Yellowdog Peridotite hand sample.
Thin sections EA-15-001, EA-15-002, EA-15-003 and EAUG0012-86.43m are from the Yellowdog Peridotite. Olivine occurs here as rounded grains, and constitutes approximately 40-60% of the unit. After olivine, pyroxene is the next most abundant phase, constituting approximately 15-40%. Most samples contain a fraction of plagioclase ranging between 15-25% of the unit. Minor minerals, such as hornblende, biotite, and magnetite account for a small percentage of the unit (<5%). Samples taken from sulfide poor sections of the unit contain minor fractions of pyrrhotite, pentlandite, and chalcopyrite. These minerals occur either as inclusions within olivines, or as small crystals filling interstitial spaces. This unit displays a cumulate texture. Olivines are subhedral to euhedral, with interstitial spaces being filled by subhedral to anhedral grains of either pyroxene or plagioclase. Poikilitic texture, in which pyroxene oikocrysts enclose olivine chadacrysts, can occasionally be observed. One example of this textural
characteristic is displayed in Figure 30. Here a group of olivine chadacrysts can be observed, completely enclosed by a clinopyroxene oikocryst. Plagioclase occurs as anhedral to subhedral laths filling the remaining interstitial space. This suggests that it was the last phase to crystallize.

Figure 29
Showing olivine, and pyroxene with interstitial plagioclase.
Geochemistry

Bulk major and minor trace element data of selected samples from each study area are presented in tables 1, and 3. Table 2 displays base metal concentrations for selected samples from each site. Tables 4, 5, 6, 7, 8, 9, 10, 11, and 12 display electron microprobe data for selected samples from Presque Isle Peridotite, Yellowdog Peridotite, and Black Rock Point Gabbro. Major oxide geochemistry numbers are expressed as weight percent. Trace element and base metal concentrations are expressed in ppm.

Figure 30
### X-ray Fluorescence Spectrometry

#### Table 6: Whole Rock Geochemistry

<table>
<thead>
<tr>
<th>Sample wt. %</th>
<th>Detection Limit</th>
<th>MQT-15-002</th>
<th>BRP-14-005</th>
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Table 7 displays base metal concentrations obtained via XRF for samples of Presque Isle Peridotite (MQT-15-002), Black Rock Point Gabbro (BRP-14-005), Eagle East (EA-15-001), and Deer Lake Peridotite (DLP-1784). Note: Elevated nickel concentrations present in peridotite samples from Presque Isle, Eagle East, and Deer Lake. This enrichment may result either from the olivine chemistry of the unit or by inclusion of nickel rich sulfide minerals. Presque Isle Peridotite also appears enriched in copper when compared to the other three units.

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Values expressed in ppm.
### Inductively Coupled Plasma Mass Spectrometry

#### Table 8: Trace Element Geochemistry

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Values expressed in ppm.
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Values expressed in ppm.
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Values expressed in ppm.
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Values expressed in ppm.
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Values expressed in ppm.
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Values expressed in ppm.
Electron Microprobe

Presque Isle Peridotite

Table 9: MQT-14-004 Olivine. Oxides expressed in wt. %

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<th>Mineral</th>
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<th>OL2</th>
<th>OL3</th>
<th>OL4</th>
<th>OL5</th>
<th>OL6</th>
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<td>290</td>
<td>293</td>
<td>294</td>
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<tr>
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<td>43.41</td>
<td>43.77</td>
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<td>0.39</td>
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<td>0.26</td>
<td>0.35</td>
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<tr>
<td>CaO</td>
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<td>0.17</td>
<td>0.10</td>
<td>0.18</td>
<td>0.20</td>
<td>0.20</td>
</tr>
<tr>
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<td>99.95</td>
<td>99.72</td>
<td>99.63</td>
<td>99.76</td>
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</table>

Stoichiometry normalized to 4 oxygens.

- Si: 1.00
- Fe(II): 0.35
- Mn: 0.01
- Mg: 1.64
- Ni: 0.01
- Ca: 0.00
- Total: 3.01

Mg# = 82.32
Fo = 82.10
Fa = 17.63

"Point" refers to analysis number or probe position.
Mg-number = 100(MgO/(MgO+FeO)) molar%
### Table 9: Continued

<table>
<thead>
<tr>
<th>Mineral</th>
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<th>OL9</th>
<th>OL10</th>
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Stoichiometry normalized to 4 oxygens.

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"Point" refers to analysis number or probe position.

Mg-number = 100(MgO/(MgO+FeO)) molar%
Table 10: MQT-14-004 Pyroxene. Oxides expressed in wt.%

<table>
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<tr>
<th>Mineral</th>
<th>PX1</th>
<th>PX2</th>
<th>PX3</th>
<th>PX6</th>
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Stoichiometry normalized to 6 oxygens.

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<th>Cr</th>
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<td>0.00</td>
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</table>

Wo 38.38 38.25 40.42 35.67 3.85 38.52 38.61
En 51.33 51.33 43.23 52.13 80.96 50.47 50.27
Fs 9.39 9.40 14.74 11.16 15.05 9.81 8.98
Ac 0.90 1.02 1.60 1.04 0.15 1.20 2.13

"Point" refers to analysis number or probe position.
Table 10: Continued

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<th>PX9</th>
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<th>PX10</th>
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<td>0.04</td>
<td>0.07</td>
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</tr>
<tr>
<td>Total</td>
<td>100.31</td>
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<td>100.19</td>
<td>100.01</td>
<td>99.54</td>
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</table>

Stoichiometry normalized to 6 oxygens.

<table>
<thead>
<tr>
<th></th>
<th>Si</th>
<th>Ti</th>
<th>Al</th>
<th>Fe(III)</th>
<th>Cr</th>
<th>Fe(II)</th>
<th>Mn</th>
<th>Mg</th>
<th>Ca</th>
<th>Na</th>
<th>K</th>
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<td></td>
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<td>0.02</td>
<td>0.02</td>
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<td>0.72</td>
<td>0.02</td>
<td>0.00</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wo</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>En</td>
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<td>1.54</td>
<td>1.19</td>
<td>1.17</td>
</tr>
</tbody>
</table>

"Point" refers to analysis number or probe position.
**Yellowdog Peridotite**

Table 11: EA-15-002 Olivine. Oxides expressed in wt. %

<table>
<thead>
<tr>
<th>Mineral</th>
<th>OL1</th>
<th>OL2</th>
<th>OL4</th>
<th>OL5</th>
<th>OL6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Point</td>
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<td>539</td>
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<td>38.76</td>
</tr>
<tr>
<td>FeO</td>
<td>18.92</td>
<td>19.09</td>
<td>18.05</td>
<td>18.29</td>
<td>18.94</td>
</tr>
<tr>
<td>MnO</td>
<td>0.18</td>
<td>0.28</td>
<td>0.26</td>
<td>0.29</td>
<td>0.20</td>
</tr>
<tr>
<td>MgO</td>
<td>41.54</td>
<td>41.19</td>
<td>41.98</td>
<td>42.06</td>
<td>41.60</td>
</tr>
<tr>
<td>NiO</td>
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<td>0.26</td>
<td>0.26</td>
<td>0.33</td>
<td>0.32</td>
</tr>
<tr>
<td>CaO</td>
<td>0.20</td>
<td>0.17</td>
<td>0.18</td>
<td>0.21</td>
<td>0.16</td>
</tr>
<tr>
<td>Total</td>
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<td>100.01</td>
<td>99.92</td>
<td>100.15</td>
<td>99.98</td>
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</table>

Stoichiometry normalized to 4 oxygens.

<table>
<thead>
<tr>
<th></th>
<th>Si</th>
<th>Fe(II)</th>
<th>Mn</th>
<th>Mg</th>
<th>Ni</th>
<th>Ca</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.997</td>
<td>0.406</td>
<td>0.004</td>
<td>1.590</td>
<td>0.005</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
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<td>1.576</td>
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<tr>
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<tr>
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<td>1.603</td>
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<td>0.006</td>
</tr>
<tr>
<td></td>
<td>0.996</td>
<td>0.407</td>
<td>0.006</td>
<td>1.592</td>
<td>0.007</td>
<td>0.007</td>
</tr>
<tr>
<td></td>
<td>0.996</td>
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<td>0.006</td>
<td>1.584</td>
<td>0.008</td>
<td>0.008</td>
</tr>
<tr>
<td></td>
<td>0.982</td>
<td>0.404</td>
<td>0.005</td>
<td>1.623</td>
<td>0.006</td>
<td>0.005</td>
</tr>
<tr>
<td></td>
<td>0.996</td>
<td>0.404</td>
<td>0.005</td>
<td>1.592</td>
<td>0.005</td>
<td>0.005</td>
</tr>
<tr>
<td></td>
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<td>0.408</td>
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<td>1.591</td>
<td>0.006</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
<td>0.996</td>
<td>0.410</td>
<td>0.007</td>
<td>1.607</td>
<td>0.006</td>
<td>0.006</td>
</tr>
<tr>
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<td>0.986</td>
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<td>1.607</td>
<td>0.006</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
<td>0.986</td>
<td>0.410</td>
<td>0.005</td>
<td>1.607</td>
<td>0.006</td>
<td>0.006</td>
</tr>
</tbody>
</table>

|   | 3.009 | 3.003 | 3.004 | 3.009 | 3.009 | 3.009 |
|   | 3.009 | 3.004 | 3.009 | 3.009 | 3.009 | 3.009 |
|   | 3.009 | 3.004 | 3.009 | 3.009 | 3.009 | 3.009 |

Mg# 79.65 79.36 80.56 80.39 79.65 79.38 80.07 79.61 79.72 79.68
Fo 79.50 79.13 80.34 80.14 79.48 79.20 79.93 79.47 79.52 79.50

"Point" refers to analysis number or probe position.

Mg-number = 100(MgO/(MgO+FeO)) molar%.
<table>
<thead>
<tr>
<th>Mineral</th>
<th>OL7</th>
<th>OL8</th>
<th>OL9</th>
<th>OL10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Point</td>
<td>552</td>
<td>554</td>
<td>560</td>
<td>561</td>
</tr>
<tr>
<td>SiO₂</td>
<td>38.51</td>
<td>38.92</td>
<td>38.68</td>
<td>38.80</td>
</tr>
<tr>
<td>FeO</td>
<td>18.58</td>
<td>18.83</td>
<td>19.18</td>
<td>18.96</td>
</tr>
<tr>
<td>MnO</td>
<td>0.21</td>
<td>0.20</td>
<td>0.20</td>
<td>0.17</td>
</tr>
<tr>
<td>MgO</td>
<td>42.13</td>
<td>41.55</td>
<td>41.49</td>
<td>41.39</td>
</tr>
<tr>
<td>NiO</td>
<td>0.22</td>
<td>0.29</td>
<td>0.25</td>
<td>0.22</td>
</tr>
<tr>
<td>CaO</td>
<td>0.19</td>
<td>0.17</td>
<td>0.17</td>
<td>0.22</td>
</tr>
<tr>
<td>Total</td>
<td>99.84</td>
<td>99.95</td>
<td>99.97</td>
<td>99.75</td>
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</table>

Stoichiometry normalized to 4 oxygens.

<table>
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<tr>
<th></th>
<th>0.989</th>
<th>0.998</th>
<th>0.994</th>
<th>0.998</th>
<th>0.996</th>
<th>0.998</th>
<th>0.993</th>
<th>0.994</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
<td>0.399</td>
<td>0.404</td>
<td>0.412</td>
<td>0.408</td>
<td>0.407</td>
<td>0.413</td>
<td>0.420</td>
<td>0.415</td>
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<tr>
<td>Fe(II)</td>
<td>0.005</td>
<td>0.004</td>
<td>0.004</td>
<td>0.004</td>
<td>0.003</td>
<td>0.004</td>
<td>0.003</td>
<td>0.003</td>
</tr>
<tr>
<td>Mn</td>
<td>1.613</td>
<td>1.589</td>
<td>1.590</td>
<td>1.587</td>
<td>1.591</td>
<td>1.580</td>
<td>1.584</td>
<td>1.587</td>
</tr>
<tr>
<td>Mg</td>
<td>0.005</td>
<td>0.006</td>
<td>0.005</td>
<td>0.004</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
</tr>
<tr>
<td>Ni</td>
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<td>0.005</td>
<td>0.005</td>
<td>0.006</td>
<td>0.003</td>
<td>0.003</td>
<td>0.004</td>
<td>0.004</td>
</tr>
<tr>
<td>Total</td>
<td>79.65</td>
<td>79.36</td>
<td>80.56</td>
<td>80.39</td>
<td>79.65</td>
<td>79.38</td>
<td>80.07</td>
<td>79.61</td>
</tr>
<tr>
<td>Mg#</td>
<td>79.98</td>
<td>79.56</td>
<td>79.24</td>
<td>79.41</td>
<td>79.49</td>
<td>79.12</td>
<td>78.91</td>
<td>79.16</td>
</tr>
<tr>
<td>Fo</td>
<td>19.79</td>
<td>20.23</td>
<td>20.54</td>
<td>20.40</td>
<td>20.34</td>
<td>20.70</td>
<td>20.91</td>
<td>20.68</td>
</tr>
</tbody>
</table>

"Point" refers to analysis number or probe position.

Mg-number = 100(MgO/(MgO+FeO)) molar%.
Table 12: EA-15-002 Pyroxene. Oxides expressed in wt. %

<table>
<thead>
<tr>
<th>Mineral</th>
<th>PX2</th>
<th>PX5</th>
<th>PX8</th>
<th>PX9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Point</td>
<td>485</td>
<td>497</td>
<td>513</td>
<td>514</td>
</tr>
<tr>
<td>SiO₂</td>
<td>53.78</td>
<td>52.09</td>
<td>51.67</td>
<td>51.42</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.54</td>
<td>0.10</td>
<td>1.12</td>
<td>0.91</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>2.07</td>
<td>29.63</td>
<td>2.66</td>
<td>3.33</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.34</td>
<td>0.00</td>
<td>0.45</td>
<td>0.74</td>
</tr>
<tr>
<td>FeO</td>
<td>11.78</td>
<td>0.74</td>
<td>6.53</td>
<td>6.55</td>
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<tr>
<td>MnO</td>
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<td>0.11</td>
<td>0.14</td>
</tr>
<tr>
<td>MgO</td>
<td>30.10</td>
<td>0.07</td>
<td>17.44</td>
<td>17.45</td>
</tr>
<tr>
<td>CaO</td>
<td>2.04</td>
<td>12.92</td>
<td>19.34</td>
<td>19.15</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.01</td>
<td>3.90</td>
<td>0.31</td>
<td>0.28</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.00</td>
<td>0.18</td>
<td>0.02</td>
<td>0.01</td>
</tr>
<tr>
<td>NiO</td>
<td>0.09</td>
<td>0.00</td>
<td>0.00</td>
<td>0.02</td>
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<tr>
<td>Total</td>
<td>100.88</td>
<td>99.56</td>
<td>99.65</td>
<td>100.01</td>
</tr>
</tbody>
</table>

Stoichiometry normalized to 6 oxygens.

Si  1.904  1.784  1.904  1.888  1.936  1.923
Ti  0.014  0.003  0.031  0.025  0.014  0.013
Al  0.000  0.980  0.019  0.033  0.000  0.000
Fe(III)  0.101  0.000  0.037  0.043  0.081  0.119
Cr  0.010  0.000  0.013  0.021  0.004  0.006
Fe(II)  0.245  0.023  0.164  0.158  0.265  0.221
Mn  0.004  0.000  0.003  0.004  0.005  0.007
Mg  1.589  0.003  0.958  0.955  1.603  1.632
Ca  0.077  0.474  0.763  0.754  0.068  0.065
Na  0.001  0.259  0.022  0.020  0.003  0.004
K   0.000  0.008  0.001  0.001  0.001  0.000
Total  3.944  3.534  3.915  3.902  3.979  3.991

Wo  3.84  62.46  39.20  38.97  3.35  3.19
En  78.79  0.45  49.19  49.40  79.18  79.68
Fs  17.34  2.97  10.47  10.58  17.32  16.95
Ac  0.04  34.11  1.13  1.05  0.15  0.19

"Point" refers to analysis number or probe position.
Table 13: EAUG0012-86.43m Olivine

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Point</th>
<th>OL4</th>
<th>OL6</th>
</tr>
</thead>
<tbody>
<tr>
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<td>621</td>
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<td>630</td>
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<td>39.31</td>
</tr>
<tr>
<td>FeO</td>
<td>17.88</td>
<td>17.45</td>
<td>16.70</td>
</tr>
<tr>
<td>MnO</td>
<td>0.16</td>
<td>0.11</td>
<td>0.12</td>
</tr>
<tr>
<td>MgO</td>
<td>42.87</td>
<td>42.93</td>
<td>43.16</td>
</tr>
<tr>
<td>NiO</td>
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<td>0.35</td>
<td>0.25</td>
</tr>
<tr>
<td>CaO</td>
<td>0.18</td>
<td>0.14</td>
<td>0.17</td>
</tr>
<tr>
<td>Total</td>
<td>100.21</td>
<td>99.77</td>
<td>99.71</td>
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</table>

Stoichiometry normalized to 4 oxygens.

<table>
<thead>
<tr>
<th>Element</th>
<th>OL4</th>
<th>OL6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
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<td>0.99</td>
</tr>
<tr>
<td>Fe(II)</td>
<td>0.38</td>
<td>0.37</td>
</tr>
<tr>
<td>Mn</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Mg</td>
<td>1.63</td>
<td>1.64</td>
</tr>
<tr>
<td>Ni</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Ca</td>
<td>0.01</td>
<td>0.00</td>
</tr>
<tr>
<td>Total</td>
<td>3.01</td>
<td>3.01</td>
</tr>
</tbody>
</table>

| Mg#     | 81.03 | 81.43 | 82.17 | 81.54 |
| Fo      | 80.90 | 81.34 | 82.06 | 81.44 |
| Fa      | 18.93 | 18.54 | 17.81 | 18.43 |

"Point" refers to analysis number or probe position.

Mg-number = 100(MgO/(MgO+FeO)) molar%
Table 14: EAUG0012-86.43m Pyroxene. Oxides expressed in wt. %

<table>
<thead>
<tr>
<th>Mineral</th>
<th>PX2</th>
<th>PX3</th>
<th>PX4</th>
<th>PX5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Point</td>
<td>647</td>
<td>649</td>
<td>651</td>
</tr>
<tr>
<td>SiO₂</td>
<td>52.63</td>
<td>52.99</td>
<td>50.49</td>
<td>50.36</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.49</td>
<td>0.59</td>
<td>1.50</td>
<td>1.27</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>2.10</td>
<td>2.26</td>
<td>3.37</td>
<td>3.45</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.55</td>
<td>0.77</td>
<td>0.27</td>
<td>0.27</td>
</tr>
<tr>
<td>FeO</td>
<td>7.47</td>
<td>6.92</td>
<td>8.74</td>
<td>8.42</td>
</tr>
<tr>
<td>MnO</td>
<td>0.08</td>
<td>0.13</td>
<td>0.17</td>
<td>0.09</td>
</tr>
<tr>
<td>MgO</td>
<td>19.37</td>
<td>18.69</td>
<td>16.57</td>
<td>16.94</td>
</tr>
<tr>
<td>CaO</td>
<td>16.94</td>
<td>18.28</td>
<td>18.43</td>
<td>18.34</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.28</td>
<td>0.23</td>
<td>0.31</td>
<td>0.31</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>NiO</td>
<td>0.04</td>
<td>0.02</td>
<td>0.06</td>
<td>0.03</td>
</tr>
<tr>
<td>Totals</td>
<td>99.93</td>
<td>100.86</td>
<td>99.90</td>
<td>99.49</td>
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</tbody>
</table>

Stoichiometry normalized to 6 oxygens.

<table>
<thead>
<tr>
<th>Element</th>
<th>PX2</th>
<th>PX3</th>
<th>PX4</th>
<th>PX5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
<td>1.93</td>
<td>1.92</td>
<td>1.87</td>
<td>1.87</td>
</tr>
<tr>
<td>Ti</td>
<td>0.01</td>
<td>0.02</td>
<td>0.04</td>
<td>0.04</td>
</tr>
<tr>
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<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
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<tr>
<td>Fe(III)</td>
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<td>0.03</td>
<td>0.05</td>
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<tr>
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<td>0.02</td>
<td>0.02</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Fe(II)</td>
<td>0.18</td>
<td>0.18</td>
<td>0.22</td>
<td>0.19</td>
</tr>
<tr>
<td>Mn</td>
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<td>0.00</td>
<td>0.01</td>
<td>0.00</td>
</tr>
<tr>
<td>Mg</td>
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<td>1.01</td>
<td>0.92</td>
<td>0.94</td>
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<tr>
<td>Ca</td>
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<td>0.73</td>
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<tr>
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<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
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<tr>
<td>K</td>
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<td>0.00</td>
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<th>PX4</th>
<th>PX5</th>
</tr>
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<td>37.64</td>
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</tr>
<tr>
<td>En</td>
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</tr>
<tr>
<td>Ac</td>
<td>1.00</td>
<td>0.85</td>
<td>1.13</td>
<td>1.13</td>
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</table>

"Point" refers to analysis number or probe position.
## Black Rock Point Gabbro

Table 15: BRP-14-006 Pyroxene. Oxides expressed in wt. %

<table>
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<tr>
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<td>52.57</td>
</tr>
<tr>
<td>TiO₂</td>
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</tr>
<tr>
<td>Al₂O₃</td>
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<td>2.30</td>
<td>2.45</td>
</tr>
<tr>
<td>Cr₂O₃</td>
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<td>0.03</td>
<td>0.28</td>
</tr>
<tr>
<td>FeO</td>
<td>9.22</td>
<td>14.32</td>
<td>10.54</td>
</tr>
<tr>
<td>MnO</td>
<td>0.30</td>
<td>0.23</td>
<td>0.29</td>
</tr>
<tr>
<td>MgO</td>
<td>15.27</td>
<td>14.04</td>
<td>17.02</td>
</tr>
<tr>
<td>CaO</td>
<td>19.66</td>
<td>16.94</td>
<td>16.86</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.22</td>
<td>0.26</td>
<td>0.23</td>
</tr>
<tr>
<td>K₂O</td>
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<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>NiO</td>
<td>0.00</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Total</td>
<td>99.58</td>
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<td>100.51</td>
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</table>

Stoichiometry normalized to 6 oxygens.

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<th>PX3</th>
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<td>6.00</td>
<td>6.00</td>
</tr>
<tr>
<td>Si</td>
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<tr>
<td>Al</td>
<td>0.04</td>
<td>0.03</td>
<td>0.04</td>
</tr>
<tr>
<td>Fe(III)</td>
<td>0.01</td>
<td>0.03</td>
<td>0.02</td>
</tr>
<tr>
<td>Cr</td>
<td>0.01</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>Fe(II)</td>
<td>0.28</td>
<td>0.42</td>
<td>0.30</td>
</tr>
<tr>
<td>Mn</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Mg</td>
<td>0.85</td>
<td>0.79</td>
<td>0.93</td>
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<tr>
<td>Ca</td>
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<td>0.69</td>
<td>0.67</td>
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<tr>
<td>K</td>
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<td>0.00</td>
<td>0.00</td>
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<tr>
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<td>4.01</td>
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</table>

<table>
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<tr>
<th></th>
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<th>PX3</th>
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<tr>
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</tr>
<tr>
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</tr>
<tr>
<td>Fs</td>
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<td>23.47</td>
<td>17.08</td>
</tr>
<tr>
<td>Ac</td>
<td>0.81</td>
<td>0.96</td>
<td>0.85</td>
</tr>
</tbody>
</table>

"Point" refers to analysis number or probe position.
Table 15: Continued

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<th>PX5</th>
<th>PX6</th>
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<td></td>
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<td>242</td>
<td>247</td>
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<tr>
<td>SiO₂</td>
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<td>51.87</td>
</tr>
<tr>
<td>TiO₂</td>
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<td>0.31</td>
<td>0.25</td>
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<td>2.99</td>
<td>2.73</td>
</tr>
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<td>Cr₂O₃</td>
<td>0.31</td>
<td>0.22</td>
<td>0.34</td>
</tr>
<tr>
<td>FeO</td>
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<td>10.02</td>
<td>9.74</td>
</tr>
<tr>
<td>MnO</td>
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<td>0.29</td>
<td>0.19</td>
</tr>
<tr>
<td>MgO</td>
<td>16.28</td>
<td>16.59</td>
<td>17.89</td>
</tr>
<tr>
<td>CaO</td>
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<td>17.58</td>
<td>16.62</td>
</tr>
<tr>
<td>Na₂O</td>
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<td>0.20</td>
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<tr>
<td>K₂O</td>
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<td>0.02</td>
<td>0.01</td>
</tr>
<tr>
<td>NiO</td>
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<td>0.01</td>
<td>0.09</td>
</tr>
<tr>
<td>Total</td>
<td>99.76</td>
<td>100.55</td>
<td>99.92</td>
</tr>
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</table>

Stoichiometry normalized to 6 oxygens.

| O  | 6.00 | 6.00 | 6.00 | 6.00 | 6.00 | 6.00|
| Si | 1.92 | 1.92 | 1.92 | 1.93 | 1.91 | 1.92|
| Ti | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | 0.02|
| Al | 0.04 | 0.05 | 0.04 | 0.04 | 0.03 | 0.03|
| Fe(III) | 0.05 | 0.03 | 0.06 | 0.03 | 0.06 | 0.05|
| Cr | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | 0.00|
| Fe(II) | 0.25 | 0.28 | 0.24 | 0.26 | 0.22 | 0.39|
| Mn | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01|
| Mg | 0.90 | 0.91 | 0.99 | 0.89 | 0.96 | 0.85|
| Ca | 0.74 | 0.69 | 0.66 | 0.75 | 0.70 | 0.65|
| Na | 0.02 | 0.02 | 0.01 | 0.01 | 0.02 | 0.02|
| K  | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00|
| Total | 4.01 | 4.01 | 4.02 | 4.01 | 4.02 | 4.01|

| Wo | 37.74 | 35.75 | 33.52 | 38.33 | 35.39 | 32.90|
| En | 46.12 | 46.97 | 50.19 | 45.54 | 48.90 | 43.34|
| Fs | 15.28 | 16.34 | 15.56 | 15.43 | 14.84 | 22.98|
| Ac | 0.86  | 0.95  | 0.72  | 0.70  | 0.87  | 0.78|

"Point" refers to analysis number or probe position.
Table 15: Continued

<table>
<thead>
<tr>
<th>Mineral</th>
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<th>PX9</th>
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<tbody>
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<td>260</td>
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<td>SiO$_2$</td>
<td>51.75</td>
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<td>51.42</td>
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<tr>
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<td>0.26</td>
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<td>0.45</td>
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<td>2.41</td>
</tr>
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<td>0.49</td>
<td>0.04</td>
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<td>8.66</td>
<td>12.80</td>
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<td>16.43</td>
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<td>0.00</td>
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<td>NiO</td>
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<td>0.07</td>
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<td>99.94</td>
<td>100.01</td>
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Stoichiometry normalized to 6 oxygens.

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<th>PX9</th>
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</thead>
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<td>6.00</td>
<td>6.00</td>
</tr>
<tr>
<td>Si</td>
<td>1.92</td>
<td>1.89</td>
<td>1.93</td>
</tr>
<tr>
<td>Ti</td>
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<td>0.01</td>
</tr>
<tr>
<td>Al</td>
<td>0.04</td>
<td>0.05</td>
<td>0.03</td>
</tr>
<tr>
<td>Fe(III)</td>
<td>0.03</td>
<td>0.06</td>
<td>0.05</td>
</tr>
<tr>
<td>Cr</td>
<td>0.01</td>
<td>0.01</td>
<td>0.00</td>
</tr>
<tr>
<td>Fe(II)</td>
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<td>Mn</td>
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<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Mg</td>
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<td>0.91</td>
<td>0.88</td>
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<td>0.66</td>
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<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>K</td>
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</tr>
<tr>
<td>Total</td>
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<td>4.02</td>
<td>4.02</td>
</tr>
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</table>

| Wo     | 36.76 | 38.52 | 33.46 | 36.71 | 36.40 | 37.83 |
| En     | 46.70 | 46.58 | 44.80 | 47.83 | 46.23 | 48.20 |
| Fs     | 15.55 | 14.00 | 20.82 | 14.51 | 16.54 | 13.16 |
| Ac     | 0.99  | 0.89  | 0.92  | 0.96  | 0.83  | 0.81  |

"Point" refers to analysis number or probe position.
Table 16: BRP-14-006 Plagioclase. Oxides expressed in wt. %

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<th>PLAG1</th>
<th>PLAG2</th>
<th>PLAG3</th>
<th>PLAG4</th>
<th>PLAG5</th>
<th>PLAG6</th>
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<td>187</td>
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<td>53.18</td>
<td>52.05</td>
</tr>
<tr>
<td>TiO₂</td>
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<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>30.33</td>
<td>29.92</td>
<td>28.98</td>
<td>30.18</td>
<td>28.89</td>
<td>29.22</td>
</tr>
<tr>
<td>Cr₂O₃</td>
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<td>0.94</td>
<td>0.91</td>
<td>1.15</td>
</tr>
<tr>
<td>FeO</td>
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<td>0.88</td>
<td>0.96</td>
<td>0.83</td>
<td>0.90</td>
<td>0.91</td>
</tr>
<tr>
<td>CaO</td>
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<td>14.02</td>
<td>12.65</td>
<td>13.21</td>
<td>12.24</td>
<td>12.28</td>
</tr>
<tr>
<td>Na₂O</td>
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<td>4.25</td>
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<tr>
<td>K₂O</td>
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<td>0.25</td>
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<tr>
<td>MgO</td>
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<td>0.00</td>
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<td>0.00</td>
</tr>
<tr>
<td>NiO</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
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<td>99.66</td>
<td>100.46</td>
<td>101.34</td>
<td>100.08</td>
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</table>

Stoichiometry normalized to 8 oxygens.

<table>
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<th>PLAG3</th>
<th>PLAG4</th>
<th>PLAG5</th>
<th>PLAG6</th>
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</thead>
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<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Al</td>
<td>6.48</td>
<td>6.41</td>
<td>6.26</td>
<td>6.24</td>
<td>6.29</td>
<td>6.46</td>
</tr>
<tr>
<td>Fe(II)</td>
<td>0.11</td>
<td>0.13</td>
<td>0.24</td>
<td>0.14</td>
<td>0.14</td>
<td>0.14</td>
</tr>
<tr>
<td>Ca</td>
<td>2.63</td>
<td>2.73</td>
<td>2.48</td>
<td>2.47</td>
<td>2.55</td>
<td>2.59</td>
</tr>
<tr>
<td>Na</td>
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<td>1.42</td>
<td>1.49</td>
<td>1.42</td>
<td>1.36</td>
</tr>
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<td>K</td>
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<td>0.04</td>
<td>0.04</td>
<td>0.05</td>
<td>0.06</td>
<td>0.03</td>
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<td>19.95</td>
<td>20.03</td>
<td>20.02</td>
<td>20.03</td>
<td>20.01</td>
</tr>
</tbody>
</table>

| An       | 67.01 | 69.24 | 62.86 | 61.57 | 63.25 | 65.15 |
| Ab       | 32.41 | 29.81 | 36.04 | 37.25 | 35.32 | 34.07 |
| Or       | 0.59  | 0.95  | 1.10  | 1.18  | 1.43  | 0.78  |

"Point" refers to analysis number or probe position.
Table 16: Continued

<table>
<thead>
<tr>
<th>Mineral</th>
<th>PLAG7</th>
<th>PLAG8</th>
<th>PLAG9</th>
<th>PLAG10</th>
<th>PLAG11</th>
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<td>216</td>
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<td>51.51</td>
<td>55.41</td>
<td>53.25</td>
<td>52.49</td>
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<td>TiO$_2$</td>
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</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>28.90</td>
<td>30.17</td>
<td>27.59</td>
<td>29.13</td>
<td>29.50</td>
</tr>
<tr>
<td>Cr$_2$O$_3$</td>
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<td>0.00</td>
<td>0.00</td>
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<tr>
<td>FeO</td>
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<td>0.61</td>
<td>1.53</td>
<td>0.86</td>
<td>0.81</td>
</tr>
<tr>
<td>CaO</td>
<td>11.37</td>
<td>13.84</td>
<td>10.41</td>
<td>12.34</td>
<td>13.15</td>
</tr>
<tr>
<td>Na$_2$O</td>
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<td>3.54</td>
<td>5.18</td>
<td>4.37</td>
<td>4.16</td>
</tr>
<tr>
<td>K$_2$O</td>
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<td>0.13</td>
<td>0.23</td>
<td>0.11</td>
</tr>
<tr>
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<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>NiO</td>
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<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Total</td>
<td>100.34</td>
<td>99.81</td>
<td>100.25</td>
<td>100.17</td>
<td>100.22</td>
</tr>
</tbody>
</table>

Stoichiometry normalized to 8 oxygens.

| Ti    | 0.00 | 0.00 | 0.00  | 0.00 | 0.00 | 0.00 | 0.00 | 0.00  | 0.00 | 0.00 |
| Al    | 6.16 | 6.49 | 5.87  | 6.23 | 6.32 | 6.47 | 5.98 | 4.21  | 6.67 | 6.55 |
| Fe(II)| 0.13 | 0.09 | 0.23  | 0.13 | 0.12 | 0.14 | 0.13 | 0.66  | 0.17 | 0.18 |
| Ca    | 2.20 | 2.71 | 2.01  | 2.40 | 2.56 | 2.68 | 2.14 | 0.15  | 2.88 | 2.84 |
| Na    | 1.76 | 1.25 | 1.81  | 1.54 | 1.47 | 1.31 | 1.83 | 3.76  | 1.15 | 1.15 |
| K     | 0.06 | 0.03 | 0.03  | 0.05 | 0.03 | 0.05 | 0.06 | 0.02  | 0.04 | 0.02 |
| Total | 20.07| 19.99| 19.97 | 20.01| 20.04| 20.04| 20.05| 20.29 | 20.08| 20.03|

| An    | 54.78| 67.85| 52.20 | 60.16| 63.18| 66.43| 53.00| 3.78  | 70.93| 70.73|
| Ab    | 43.68| 31.36| 47.01 | 38.52| 36.18| 32.41| 45.45| 95.79 | 28.17| 28.72|
| Or    | 1.54 | 0.79 | 0.79  | 1.32 | 0.64 | 1.16 | 1.54 | 0.44  | 0.90 | 0.55 |

"Point" refers to analysis number or probe position.
Data Analysis

Tables 9 through 16 display geochemical data collected via electron microprobe for samples of: (1) Presque Isle Peridotite, (2) Yellowdog Peridotite (from both the Eagle and Eagle East intrusions), and (3) Black Rock Point Gabbro. Stoichiometry has been calculated for each crystal that was probed, and normalized to the number of oxygens in the chemical formula of that mineral.

End member compositions of each probed mineral were also calculated, and are presented as a molar percentage in tables 9 through 16. This was done according to the procedure outlined by the serc.carleton.edu website by Brady and Perkens. The weight percentage of each oxide was divided by the formula weight of that oxide. Secondly, the resulting "mole number" of each oxide was multiplied by the number of oxygens in the oxide formula. The resulting "oxygen number" of each oxide was then multiplied by a normalization constant (equal to the number of oxygens in the desired formula divided by the sum of the "oxygen numbers"). Finally, the "normalized oxygen numbers" of each oxide were multiplied by the number of cations per oxygen in the oxide formula. Only then could the percentages of each end member, present in each probed crystal, be calculated according to their elemental makeup.
Figure 31 is the ternary diagram known as a “pyroxene quadrilateral”. The diagram displays the distribution of pyroxene compositions for the individual crystals probed in each sample. According to the diagram, all of the probed pyroxenes from Black Rock Point Gabbro, and the majority of those from Eagle, Eagle East, and Presque Isle Peridotites can be identified as augite (a clinopyroxene mineral). Additionally, several probe points from Eagle East, and one from Presque Isle, can be identified as enstatite (an orthopyroxene mineral).

Figure 32 shows the distribution of olivine end member compositions among three samples from East Eagle, Eagle, and Presque Isle. Olivine from all three sites shows a predominately forsteritic composition with East Eagle, or the eastern intrusion of Yellowdog Peridotite showing the highest average fayalite concentration per olivine.
crystal probed, and Presque Isle Peridotite showing the lowest. Despite the observed
differences in forsterite content, the average end member composition of probed olivines
from all three sites fall within a range of less than 4% forsterite. Given the relatively
small sampling of probe points from each site, this variation would be expected to
decrease further were the sample size to be increased.

**Olivine Composition**

- EA-15-001
- EAUG0012-86.43m
- MQT-14-003

**Figure 32**
Plot of olivine compositions for East Eagle, Eagle, and Presque Isle.
CHAPTER V
DISCUSSION

Petrology

All three peridotite units have been observed to exhibit a cumulate texture, consisting of a fabric of interlocking subhedral to euhedral olivine with interstitial spaces filled by pyroxene (or, as in the case of the Yellowdog Peridotite, pyroxene and plagioclase). These three peridotites also exhibit poikilitic texture, in which pyroxene oikocrysts, fully or partially enclose olivine chadacrysts. Both textures are common features of cumulate type peridotite units (Wagner et al., 1960). Therefore it can be assumed that these three peridotite units resulted from fractional crystallization of an intrusive magma, and are not examples of mantle peridotite that was tectonically emplaced.

All three peridotite units display different degrees of serpentinization. Deer Lake Peridotite has been almost entirely serpentinized. Presque Isle Peridotite has been partially to mostly serpentinized depending upon where the unit is sampled. Yellowdog Peridotite displays by far the lowest degree of serpentinization. Serpentinization results from the interaction of high temperature and pressure minerals (such as olivine and pyroxene, which are unstable at near surface conditions) with water and oxygen. Oxidation and hydrolysis reactions occur, resulting in alteration of primary olivine and pyroxene, into secondary serpentine. Differing degrees of alteration may be attributed to: (1) Differences in time since formation, (2) exposure to near surface conditions of temperature and pressure, and (3) variance in the number of episodes, duration, and intensity of these hydrothermal interactions.
Yellowdog Peridotite, in contrast to Marquette County’s other two peridotite units, contains a substantial fraction of plagioclase feldspar (primarily as interstitial crystals). Findings of Saper and Yang (2014) suggest that the mineralogy of this unit may result from the “refertilization” of residual peridotite with mafic melt. If the Yellowdog Peridotite is considered to have formed as a cumulate (as concluded above). Any secondary reaction of the peridotite with its remaining parental melt could explain the feldspathic composition of the unit.

In Presque Isle Peridotite, pentlandite and chalcopyrite were observed in association with magnetite. Nickel sulfide is present as the hypogene mineral, pentlandite, not its supergene equivalent, violarite, which is common in hydrothermal deposits, therefore, it is likely that these mineral assemblages are representative of primary magmatic sulfides, hosted within this peridotite unit.

Figure 33
Comparison of sulfides from Presque Isle Peridotite (A) and Yellowdog Peridotite (B).
In Yellowdog Peridotite, pyrrhotite, in addition to pentlandite, and chalcopyrite can be observed in association with magnetite. A comparison of sulfide mineral assemblages from both units can be viewed in Figure 33.

Black Rock Point Gabbro is composed primarily of augite and plagioclase feldspar. The unit also contains only a small percentage of sphene, and no olivine is present. These characteristics allow the major mafic unit found at Black Rock Point to be classified as a gabbro (per the IUGS classification scheme, shown in Figure 34).

*Figure 34*
IUGS classification scheme for gabbro. (Woolley, 1996)
Geological Mapping

Maps of each field area were created using hand drawn field maps based on data collected at each site. These maps were then used in conjunction with previously published maps, for Presque Isle by, Gair and Thaden (1968), and for Deer Lake by, Clark, Cannon, and Klasner (1975). Finally, field maps were digitized using Esri ArcMap software to produce the final maps for each area.

Presque Isle Peridotite

![Geological map of Presque Isle](image)

**Figure 35**
Geological map-Presque Isle.
Adapted from Gair and Thaden (1968).
Figure 36
Geological map-Deer Lake area.
Adapted from Clark, Cannon, and Klassner (1975).
Figure 37
Geological map-Black Rock Point showing sample locations from May 2014 and 2015.
Geochemistry

Electron Microprobe

Olivine Data collected via electron microprobe can be used to determine the MgO/(MgO+FeO) ratio of each crystal’s parental magma. The results of these calculations, averaged for each sample, appear in Table 13. According to Tatsumi et al (1983), the MgO/(MgO+FeO) ratio of a parental mantle magma can be expected to fall between 0.63 and 0.73.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Average MgO/(MgO+FeO)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Presque Isle (MQT-14-003)</td>
<td>0.72</td>
</tr>
<tr>
<td>Eagle East (EA-15-001)</td>
<td>0.69</td>
</tr>
<tr>
<td>Eagle East (EUG0012-86.43m)</td>
<td>0.71</td>
</tr>
</tbody>
</table>

Major element analysis has revealed that all three peridotite units appear to be anomalously rich in nickel (Table 7). This is not surprising in the case of Yellowdog Peridotite. The unit is already observed to host a rich Cu-Ni magmatic sulfide deposit. High nickel concentrations in the other peridotite units may also be due to nickel-rich sulfides. However, the high nickel anomalies may also result from a high concentration of nickel in contained in the crystal structure of olivines. This can be determined by using data collected from electron microprobe analysis. Figure 37 displays a plot of Ni-content vs. Forsterite molar percent. This comparison can be used to determine if nickel depletion of olivine, in the presence of an immiscible sulfide liquid, has occurred.
Maier et al. (2015) suggests that intrusions whose olivines show a steep trend from high Fo and Ni toward low Ni content, when plotted in this manner, are prospective. This trend suggests that sulfide segregation has taken place within the intrusion. Any intrusion which contains only Ni depleted olivine suggests that the nickel depletion must have occurred prior to the final emplacement of the intrusive body. There is a low probability that any such body would be able to host a magmatic sulfide deposit.

Ding et al. (2010) used a plot similar to Figure 38, to model olivine compositions from both intrusions of the Yellowdog Peridotite. Crystallization curves were also constructed based on the olivine composition of the Keweenawan, Mamainse Point Formation. This model has been applied to data collected by this study (Figure 37).

**Figure 38**
Ni-content vs. Fo mol% plot. After, Ding et al. (2010).
Blue curve: Equilibrium crystallization.
Green curve: Fractional crystallization.
Olivine compositions samples MQT-14-003, EA-15-001, and EUG0012-86.43m which were analyzed using electron microprobe, display a range of nickel contents which spread across both the fractional and equilibrium crystallization curves. Compositions of olivine ranged from high Fo/high Ni, to high Fo/low Ni. Olivine which plotted above the fractional crystallization curve shows Ni enrichment, while those which plot below show Ni depletion.

Similar trends are displayed by all three intrusions. Both Eagle and Eagle East host massive sulfide deposits. This fact, along with the similar Ni/Fo trends of the three intrusions, suggests that these intrusions have all experienced Ni depletion of olivine, in the presence of an immiscible sulfide liquid. Additionally, the findings suggest that Presque Isle Peridotite may also host Ni-Cu-PGE sulfide mineralization, and should be considered prospective. However, the small sample size utilized in cannot be relied upon as a basis for absolute conclusions.
Major and Minor Trace Elements

Figures 39, 40, and 41 compare all four rock units targeted by this study in terms of minor and trace element composition.

A high degree of variation, in terms of minor trace element composition, seems to exist between samples of Deer Lake Peridotite. When viewing only Figure 39, one might consider this pattern as the product of the extensive alteration which has occurred in the Deer Lake Peridotite. However, when only the immobile elements are considered (as shown in Figure 40) variations continue to be apparent. Type 2 Deer Lake Peridotite (sample DLP-1784) shows enrichment in Hf, Zr, and Tm which is not present in Type 1
Deer lake Peridotite (sample DLP-15-006). Geochemical variations between samples taken from the Deer Lake Peridotite may imply that the unit was not formed by a single magmatic pulse. Formation of this unit may actually have resulted from multiple episodes of magmatic intrusion.

![Spider plot - NMORB immobile (Sun & McDonough 1989 in Pearce 2014)](image)

**Figure 40**
Immobile element spider plot. Normalized to NMORB.

Due to their incompatibility, rare earth elements are particularly useful for identification of a suite of rocks formed as a result of crystal fractionation (Nelson, 2012). Such a suite should plot as a group of nearly parallel lines showing varying degrees of rare earth element enrichment. Figure 41 shows a data plot for these elements. Four distinct suites can be identified on this graph. Samples of Black Rock Point Gabbro appear to plot as one suite at the top of the graph. Samples of Type 1 and Type 2 Deer
Lake Peridotite plot as two separate suites. Samples of both Presque Isle Peridotite and Yellowdog Peridotite plot as a fourth suite of seven nearly parallel lines.

Spider plot – REE Primitive mantle (McDonough and Sun 1995)

![Spider plot](image)

Figure 41
Rare earth element spider plot. After McDonough and Sun (1995).

The suite discussed (Black Rock Point Gabbro) was entirely composed of multiple samples from a single unit; therefore, their geochemical relationship is expected. Trace element variation between Type 1 and Type 2 Der Lake Peridotite is not entirely unexpected, and may constitute additional evidence suggesting that this unit was formed by two separate magmatic events. The Nearly identical trends displayed by samples from both Eagle, and Presque isle are significant. The geochemical similarity between these units is not isolated to rare earth elements, but can also be observed in both Figure 39 and Figure 40. There is a clear geochemical relationship in terms of minor-trace elements
between these two rock units. This similarity exists despite the fact that the dates of formation for these units are conventionally thought to be nearly 1 billion years apart.

**Geotectonic Proxies**

Two geotectonic proxies were applied to the geochemical data in an effort to determine the origin of each unit. These are the Th-Nb proxy for crustal input and the Ti-Yb proxy for melting depth. This was conducted according to the methodology established by Pearce (2008). The proxies are based on the following principles which can be observed by plotting a range of samples on a MORB normalized trace element similar to Figure 38. These proxies function according to the following principles (shown visually in Figure 42).

1. Crustal contamination of a mafic melt by continental lithosphere will result in selective Th and LREE enrichment and thus negative Nb anomalies. This allows for the use of Th-Nb as a proxy for crustal input. Such an Nb

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**Figure 42**
depletion can be observed for all samples in Figure 39.

(2) Source and melting effects result in Nb-Ti gradients. Nb-Ti gradients are indicative of incompatible enrichment which cannot be explained by HREE compatibility in garnet. This gradient increases from tholeiitic to alkalic compositions.

(3) Presence of garnet residue will result in negative Ti-Yb gradient. Ti and Yb are portioned similarly in spinel peridotite, while Yb is much more strongly partitioned into garnet. This allows for Ti/Yb to be used as a proxy for melting depth.

Figure 43 and Figure 44 display ratio plots designed to highlight crustal input, and melting depth proxies by focusing on Yb normalized ratios (refer to Figure 42b). Figure 43 is an Nb/Yb-Th/Yb plot based on Pearce (2008). This diagram is a graphical representation of the Th-Nb crustal input proxy. Present day MORB and OIB form a diagonal array with average N-MORB, E-MORB, and OIB at its center. This array contains >98% of analyzed oceanic basalts. Melts which have interacted with continental crust during ascent, display higher Th/Yb values (Pearce, 2008). Samples from all four research areas plot above the diagonal array. This indicates that the melts from which they formed were subjected to crustal contamination either during their intrusion, or as the result of subduction related input prior to crystallization of each, individual unit.
Figure 43
Nb/Yb-Th/Yb plot. Ratio plot highlighting Th-Nb proxy.

Figure 44 is a TiO\textsubscript{2}/Yb-Nb/Yb plot based on Pearce (2008). This plot is a graphical representation of the Ti-Yb melting depth proxy. TiO\textsubscript{2}/Yb values which plot above the MORB array are indicative of melting which has occurred beneath thicker lithosphere. Data acquired from samples taken from each research area in this study are included on this plot. These data split into two distinct groupings. Both Deer Lake Peridotite and Black Rock Point Gabbro plot within the shallow melting array. Samples
of Yellowdog Peridotite and Presque Isle Peridotite Plot with in the deep melting zone as indicative of their elevated TiO2/Yb values.

**Figure 44**
TiO2/Yb-Nb/Yb plot. One sample from each site is represented. Ratio plot highlighting Ti-Yb proxy.

The environment of formation for each unit can be proposed when both the Th-Nb crustal input proxy and the Ti-Yb melting depth proxy are considered. Samples of both Deer Lake Peridotite and Black Rock Point Gabbro display elevated Th/Yb values indicating of crustal contamination of their parental magmas. However, neither unit displays elevated TiO2/Yb values. This suggests that both are products of shallow melting. The parental melts of these units were most likely formed as a result of shallow dipping ocean-ocean or ocean-continent subduction. Samples of Yellowdog Peridotite
and Presque Isle Peridotite display elevated Th/Yb values indicating crustal contamination. Elevated TiO$_2$/Yb indicative of deep melting is also present in both units. Evidence, in this case, suggests that parental magmas were probably formed as a result of plume-initiated melting deep beneath the continental lithosphere. When the melts intruded through the upper mantle and continental crust, they became contaminated, prior to their crystallization.

Conclusion

**Deer Lake Peridotite**

Mineralogical compositions and textural characteristics of Deer Lake Peridotite are similar to those observed in the Presque Isle Peridotite and Yellowdog Peridotite. Prior to alteration, the unit’s mineralogical composition was dominated by olivine and pyroxene. This rock type displays a cumulate texture in some cases, a poikilitic texture in which pyroxene oikocrysts enclose olivine chadacrysts is also present. However these similarities are not sufficient to conclude that this unit shares a common origin with either of the other peridotite units. Also, it is noteworthy that the degree of serpentinization and hydrothermal alteration observed in the Deer Lake Peridotite is different, being far greater than that observed in either of Marquette County’s other peridotite units.

Geochemical comparison of Deer Lake Peridotite with the other three units addressed in this study reveals obvious differences in chemical composition. Geochemical analysis also reveals that the Deer Lake Peridotite crystallized from a
magma which formed as a result of shallow melting; whereas, the parent magmas of Presque Isle Peridotite and Yellowdog Peridotite formed as a result of deep melting.

Truncation of the Deer Lake Peridotite along its south-western margin by the Great Lakes Tectonic Zone suggests that the unit must have been formed either during, or prior to the formation of the GLTZ (2.7-1.85 Ga). This window of time for this formational event proves that the Deer Lake Peridotite predates the Yellowdog Peridotite’s age (1.1 Ga) by no less than 750 Ma.

Trace element, and petrographic analysis also suggests that Deer Lake Peridotite may actually represent two separate peridotite units emplaced during two separate events. Type 2 Deer Lake Peridotite displays foliation which is not present in Type 1 Deer Lake Peridotite. Possibly, Type 2 Deer Lake Peridotite was emplaced early in the formation of the GLTZ, and later deformed during the Penokean Orogeny (1.86-1.83 Ga), with Type 1 crystallizing during this later compressive phase. Bornhorst et al. (1993) postulated that the Deer Lake Peridotite may represent the subvolcanic base of the Mona Formation. Deer Lake Peridotite also appears to be associated with the metavolcanics of the Kitchi Formation. It is possible that these successive metavolcanic units may correspond to the two ultramafic units of Deer Lake Peridotite.

Based on geochemical analysis, in conjunction with the geologic setting of the Deer Lake Peridotite, it is reasonable to conclude that the Deer Lake Peridotite was formed independently, and substantially earlier, than Presque Isle Peridotite and Yellowdog Peridotite. It is likely that this unit crystallized from a parent magma which resulted from shallow melting during the formation of the Great Lakes Tectonic Zone.
**Black Rock Point Gabbro**

Petrological comparison of Black Rock Point Gabbro with the other three units addressed by this study yielded little similarity. Unlike Marquette County’s three peridotite units, Black Rock Point Gabbro contains no olivine. The unit’s primary minerals include augite and plagioclase feldspar, with small amounts of sphene. Additionally, no shared textural characteristics were identified between the Black Rock Point Gabbro, and any of the Marquette County peridotites.

Geochemical comparison of Black Rock Point Gabbro with the Marquette County Peridotites reveals a substantially different minor trace element signature. Geochemical analysis also leads to the conclusion that the unit crystallized from a parent melt which resulted from shallow melting before becoming contaminated by crustal material. This environment of formation suggests it is likely that the Black Rock Point Gabbro also dates from the formation of the Great Lakes Tectonic Zone (2.7-1.85 Ga).

Due to the limited extent of the Black Rock Point Gabbro’s outcrop, very little can be concluded based on its geologic setting. It does intrude the Late-Archean granite of the “Northern Complex”, and it is overlain by the Jacobsville Sandstone nonconformably.

Based on petrographic analysis, no relationship can be established between the Black Rock Point Gabbro and the other three units addressed in this study. Although geochemical analysis suggests that both the Black Rock Point Gabbro and Deer Lake Peridotite crystallized from parent melts formed under similar circumstances, their
substantially different geochemical compositions give no indication that these melts were similar or directly related.

**Presque Isle Peridotite and Yellowdog Peridotite**

Presque Isle Peridotite and Yellowdog Peridotite, upon first examination, appear to be very different units. Presque Isle Peridotite is more finely grained, and lacks visible plagioclase feldspar crystals, such as those observed in the Yellowdog Peridotite. Presque Isle Peridotite has also undergone a notably higher degree of serpentinization, as is evidenced by a greater density of hydrothermal veins.

Additional examination also reveals that primary mineral assemblages in both units include a large fraction of olivine. Pyroxene, mostly in the form of augite, with a much smaller fraction of enstatite, also constitutes a substantial fraction of both units. They both display a cumulate texture. Additionally, a poikilitic texture in which rounded olivine chadacrysts are partially, or fully enclosed by pyroxene oikocrysts can be observed in both the Presque Isle Peridotite and Yellowdog Peridotite. Petrographic analysis of both units also reveals olivine hosted sulfide inclusions. Such inclusions may indicate the presence of an immiscible sulfide liquid at the time of crystallization.

Geochemical comparison of the Presque Isle Peridotite and Yellowdog Peridotite indicates that both formed from a parent melt of mantle origin, and became contaminated by crustal material prior to crystallization. Minor trace element analysis of the two units, reveals that they share a very similar geochemical composition. Samples from both sites plot as a tightly spaced group of nearly parallel lines (Figures 39, 40, and 41).
Limited outcrop exposure of the Yellowdog Peridotite, and the lack of accessibility to host rock contacts at the Presque Isle Peridotite, make it difficult to draw conclusions based strictly on geologic setting relationships at both sites. However, Yellowdog Peridotite has intruded through the Late-Archean granite basement of the “Northern Complex”, and Paleoproterozoic sediments of the Baraga Basin (as confirmed by the exploration teams of both Kennecott and Lundin Mining). Ding et al. (2010), by the use of U-Pb baddeleyite dating, has confirmed that this unit crystallized at 1107.2 ± 5.7 Ma. This date allows for the reasonable conclusion that the Yellowdog Peridotite formed during the Midcontinent Rift event. The only unit which has been observed in direct contact with the Presque Isle Peridotite is the uppermost member of the Keweenawan series, the Jacobsville Sandstone. Here, it is clear that Presque Isle Peridotite is nonconformably overlain. Radiometric dating of zircons from Jacobsville Sandstone confirms that the unit is no younger than 960 Ma (Malone et al., 2015). Observation of the peridotite’s contacts with other units is not possible because they are concealed beneath Lake Superior. However, it can be safely assumed that the Presque Isle Peridotite has also intruded through the Archean granite basement of the “Northern Complex”. No geochronological dates have ever been obtained for the Presque Isle Peridotite.

In summary, Presque Isle Peridotite and Yellowdog Peridotite share similar mineralogical and textural characteristics. Geochemical analysis reveals that both units crystallized from parent magmas of a mantle origin, which were contaminated by crustal material during their intrusion and prior to crystallization. Analysis of minor trace
elements also reveals that the units share a very similar chemical signature, and can easily be classified as a single suite of rocks.

The findings also suggest the possibility that the Presque Isle Peridotite may also date to the Mesoproterozoic, at which time, it may have been formed contemporaneously with the Yellowdog Peridotite, during the early stages of the Midcontinent Rift event (1.1 Ga). Geochemical similarities between these units also suggest the likelihood that the plume induced, parent melts of both units were very similar, and may have been directly related.

Presque Isle Peridotite is shown to display the following: (1) High nickel content, comparable to that of the Yellowdog Peridotite (as shown by XRF analysis). (2) Sulfide inclusions within olivine. (3) Primary magmatic sulfide assemblages of chalcopyrite and pentlandite. (4) Incompatible element enrichment. Comparison of Ni content with Fo molar percentage, also yields results similar to those observed in both the “Eagle” and “Eagle East” intrusions of Yellowdog Peridotite. All these factors make Presque Isle Peridotite a prime target for future exploration, as they suggest the possibility that the unit has the potential to host magmatic sulfide deposit similar to those hosted by both intrusions of the Yellowdog Peridotite. It can be concluded from the available data, that peridotite units of Keweenawan age, located in the Lake Superior region should be considered high priority targets for magmatic sulfide exploration.
Future Work

Geochronologic analysis of Presque Isle Peridotite should be the top priority. If age dates can be obtained using U-Pb baddeleyite dating, as was accomplished for the Yellowdog Peridotite, the debate over the age of the Presque Isle Peridotite can finally be put to an end. Dates returned will greatly solidify any concepts concerning the origin of the Presque Isle Peridotite.

Additional geochemical analysis should also be conducted using XRF and ICP-MS for both Presque Isle Peridotite and Yellowdog Peridotite. This new data should be combined with data developed in this study, in an effort to determine if the geochemical similarity between Presque Isle Peridotite and Yellowdog Peridotite can be further supported by a larger, more statistically valid, sample size.

Expanded electron microprobe studies of Presque Isle Peridotite should also be undertaken. Additional data from a wide range of samples will allow for improved modeling of nickel depletion currently observed in olivines of Presque Isle Peridotite. This will allow for a better assessment of this unit as a potential magmatic sulfide host.

Sulfur isotope analysis of primary magmatic sulfides from Presque Isle Peridotite and Yellowdog Peridotite should be conducted. This analysis would allow for the origin of sulfur to be determined for these sites.

Metallic ore bodies are associated with significant magnetic anomalies. Aeromagnetic surveys played a role in the discovery of the Eagle magmatic sulfide deposit, hosted within the Yellowdog Peridotite, and have continued to be employed by subsequent exploration efforts in the area. Aeromagnetic survey data which includes the
Presque Isle area has never been made publically available. Analysis of Presque Isle Peridotite’s associated magnetic anomaly would not only help to define the most promising locations for magmatic sulfide exploration, but would also reveal the full extent of the unit, which is concealed beneath Lake Superior.
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