Peridotite melting and mineral–melt partitioning of major and minor elements at 22–24.5 GPa

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Abstract

Melting experiments at 2100–2400°C and 22–24.5 GPa to determine the liquidus to solidus phase relations and element partitioning coefficients were performed on different peridotite compositions: a pyrolite model and a KLB-1 composition. These starting materials, prepared as oxide mixes, were contained in Re capsules. The experiments were carried out using cubic-octahedral multianvil presses with 10/4 mm (octahedral edge length/anvil truncation edge length) and 18/8 mm pressure cell configurations with LaCrO3 heaters and axially inserted W-Re thermocouples. The 18/8 mm configuration with a stepped heater was developed in order to reduce the experimental thermal gradients. Some of the experiments were performed with Re-rod sample containers with two parallel chambers, allowing two different compositions to be run at identical conditions. Within experimental uncertainties, the melting relations of the KLB-1 and pyrolite compositions are identical in the 22–24.5 GPa range. Ringwoodite does not occur either in the melting range or just below the solidus, even at 22 GPa. Garnet (ga) and ferropericlase (fp) are the first liquidus phases at 22–23 GPa and at 24–24.5 GPa, respectively. The crystallization sequence is ga-fp-Ca-perovskite (cpv) at 22 GPa, ga-fp-perovskite(pv)-cpv at 23 GPa, and fp-pv-ga-cpv at 24–24.5 GPa. Major element distribution between the different phases implies that the solidus is approximately coincident with the appearance of cpv. The major element mineral–melt partitioning shows only minor variations with pressure in the investigated range. The mineral–melt $K_{D}(\text{Fe/Mg}) = (\text{Fe/Mg})_{\text{mineral}}/(\text{Fe/Mg})_{\text{melt}}$ decreases from 0.64 at 22–23 GPa to 0.62 at 24.5 GPa for ferropericlase and is below 0.46 for garnet and perovskite. The $K_{D}(\text{Si/Mg})$ for garnet and perovskite decrease slightly from 1.34 at 22 GPa to 1.16 at 24.5 GPa. The Ca/Al ratio is elevated in the melt phase with mineral–melt $K_{D}(\text{Ca/Al})$ for garnet and perovskite below 0.4. The partitioning coefficients indicate that partial melting or fractional crystallization in the transition zone and the uppermost part of the lower mantle will produce melts with elevated ratios of Fe/Mg and Ca/Al. The partitioning of minor elements like Ti, Cr, Ni and Na is strongly dependent on the proportions of the fractionating phases. © 2002 Elsevier Science B.V. All rights reserved.

Keywords: peridotites; high pressure; melting; perovskite; garnet group; periclase

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1. Introduction

Partial melting of mantle peridotite at pressures below 5 GPa produces the vast majority of mafic magma types seen at the Earth’s surface. At various times during the early history of the Earth, however, peridotite melting may have occurred at much higher pressures, corresponding to the transition zone and the upper part of the lower mantle. Impacts of giant planetesimals during the later stages of the accretion of the Earth may have resulted in partial or complete melting of the mantle [1]. Fractional crystallization of such a magma ocean may have produced an early stratification of the mantle, although convective stirring may have greatly reduced such a layering [2]. Hotter plumes rising through the Archean mantle, however, may also have experienced initial melting in the upper part of the lower mantle [3,4].

The lack of precise and reliable element partitioning coefficients between melt and residual minerals stable in the lower mantle and the transition zone has caused uncertainties and debate over the possible consequences of fractionation processes related to the crystallization of a magma ocean and to melt separation from hot plumes (e.g. [5,6]). A knowledge of the melting phase relations and element partitioning expected during melting of peridotite at high pressure is therefore of great interest. In addition, the study of equilibrium partitioning between phases at high pressure is often hindered by the difficulty in attaining equilibrium at plausible mantle temperatures. During melting, however, rapid transport in the liquid aids the attainment of equilibrium and crystal-chemical influences over element partitioning between solids and between solid and liquid phases can be studied.

Although several studies have examined melting relations of peridotitic and chondritic compositions and mineral-melt partitioning at transition zone and lower mantle conditions (e.g. [7–14]), the solidus to liquidus phase relations above 22 GPa are still not well established. The mineral-melt partitioning of major and trace elements above 20 GPa, including the effect of pressure, is also uncertain. Here we report results of melting experiments performed on mantle peridotite compositions between 22 and 24.5 GPa. These new data were collected with specific attention paid to the minimization of thermal gradients at high pressure. A selection of 20 trace elements doped at the 60–800 ppm level (in total 0.53 wt%) were added to some of the starting powders in order to extract trace element partitioning data, which will be reported separately.

2. Experimental technique

Table 1 lists the experimental conditions and observed phase relation. The major element compositions of the KLB-1 and pyrolite starting compositions are given in Table 2. The starting compositions were produced from reagent grade oxides and carbonates, which were ground together in alcohol and decarbonated at 1000°C in air. A selection of 20 trace elements with an oxide total of 0.61 wt% were added to the pyrolite composition using AAS standard solutions. The powdered mixes were subsequently devolatilized at 400°C in air. Cold pressed pellets of starting powders were reduced in a CO2-CO gas mix furnace at 1200°C and at an fO2 of two log units below the fayalite-magnetite-quartz oxygen buffer. The recovered pellets were reground to a grain size of 5–10 μm.

In order to minimize thermal gradients during the experiments, sample capsules were kept as short as possible. Re foil capsules 1 mm long were used and in addition multichamber capsules were fabricated from 0.6 mm long Re rods by spark-eroding the sample holes. Two sample chambers, 0.3 mm in diameter, in each rod allowed two starting materials to be run in a single experiment.

Experiments were conducted in a 1200 ton split sphere multianvil apparatus using a 10 mm edge length Cr2O3-doped MgO octahedron and 32 mm WC cubes with 4 mm edge length truncations. In order to investigate the melting relations in an environment of more reduced thermal gradients, some of the experiments were performed with larger pressure cells (18 mm octahedral edge lengths) in a 5000 ton split cylinder multianvil press, using 54 mm Toshiba F grade cubes with
8 mm edge length corner truncations. In the latter configuration, a pressure of 23 GPa was reached by press loads of 2000 tons. The larger 18 mm octahedra provide space for narrow and stepped LaCrO₃ furnaces that resulted in thermal gradients (determined by two-pyroxene thermometry), on the order of 50°C/mm at 1400°C, as opposed to gradients of more than 100°C/mm in a 10 mm octahedron at the same temperature.

The high-temperature pressure calibrations were performed at 1600°C and 2000°C. The calibration curves are based on the following transitions in the compositions MgSiO₃ and Mg₂SiO₄: forsterite–wadsleyite, wadsleyite–ringwoodite and ilmenite–perovskite [15,16], with reference to the Jamieson et al. [17] equation of state for gold. An additional calibration check point at 26 GPa was provided by the compositions of coexisting garnet and perovskite in the enstatite–pyrope system [18]. The pressure calibration has been adjusted relative to preliminary reports [19,20]. Using the recently modified pressure scale [21,22] based on the Anderson et al. [23] equation of state for gold, our pressure values must be lowered by about 2 GPa. The current uncertainties [24,25] and the advantage of direct comparison with the pressure

### Table 1
List of experiments

<table>
<thead>
<tr>
<th>Run number</th>
<th>Starting material</th>
<th>p (GPa)</th>
<th>Nominal T (°C)</th>
<th>Duration (min)</th>
<th>Mineral assemblage (crystallization sequence)</th>
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<tr>
<td>2377</td>
<td>py</td>
<td>22</td>
<td>2150</td>
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<td>py*</td>
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<td>2160</td>
<td>20</td>
<td>L-ga-fp-cpv</td>
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<tr>
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<td>py</td>
<td>22</td>
<td>2160</td>
<td>20</td>
<td>L-ga-fp-cpv</td>
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<td>22</td>
<td>2250</td>
<td>20</td>
<td>L-ga-fp</td>
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<td>23</td>
<td>2140</td>
<td>20</td>
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<td>23</td>
<td>2160</td>
<td>15</td>
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<tr>
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<td>–</td>
<td>15</td>
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<tr>
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<td></td>
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<td>2230</td>
<td>40</td>
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<td>2270</td>
<td>10</td>
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<tr>
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<td>L-fp-pv-ga</td>
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<tr>
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<td>24.5</td>
<td>2270</td>
<td>20</td>
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<tr>
<td>KLB</td>
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<td></td>
<td>L-fp-pv-ga</td>
</tr>
<tr>
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<td>py</td>
<td>24.5</td>
<td>–</td>
<td>3</td>
<td>above liquidus</td>
</tr>
<tr>
<td>KLB</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>L-fp</td>
</tr>
<tr>
<td>2425</td>
<td>py</td>
<td>24.5</td>
<td>2400</td>
<td>2</td>
<td>L-fp</td>
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</table>

18–8 mm assemblies were used for experiments at 23 GPa, 10–4 mm assemblies at all other pressures. Sample capsules: Re-foil, except for experiments 2276 and 2383 (Re-rod capsules with one sample chamber) and all 24.5 GPa experiments (Re-rod capsules with two sample chambers). The nominal temperature was recorded by the thermocouple during heating. After arrival at run temperature, the experiments were kept at constant electrical power output. Temperatures are not given for two experiments with no thermocouple control during heating.

Starting materials: py, pyrolite [29] with added trace elements; py*, pyrolite without trace elements; py+Fe, pyrolite with trace elements and an extra addition of 1% Fe metal; KLB, KLB-1 peridotite [30] without trace elements. Mineral assemblages: L, melt; iL, inferred interstitial melt; ga, majoritic garnet; fp, ferropericlase; pv, MgSi-perovskite; cpv, CaSi-perovskite.
values of published phase equilibrium studies on peridotite compositions make the use of the old pressure scale \([15^\text{17}]\) our natural choice. The temperature was measured using WRe3/WRe25 thermocouples with no pressure correction of the thermocouple EMF.

The individual components of the pressure assembly, including octahedra, ZrO\(_2\) insulating sleeves, MgO spacers and the LaCrO\(_3\) heater parts, were fired at 1000°C for 1–2 h. After complete assembly the octahedra with capsules were kept at 250°C in a vacuum oven for at least 10 h before an experiment. Experiments were performed by increasing the pressure over a duration of 4 h, followed by slow heating to 1400°C (250°C/min). The temperature was kept at 1400°C for 20 min, before rapid heating to the run temperature (500°C/min). The nearly linear

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### Table 2
Composition of starting materials, quenched melts and coexisting liquidus minerals

<table>
<thead>
<tr>
<th></th>
<th>SiO(_2)</th>
<th>Al(_2)O(_3)</th>
<th>TiO(_2)</th>
<th>Cr(_2)O(_3)</th>
<th>NiO</th>
<th>FeO</th>
<th>MnO</th>
<th>MgO</th>
<th>CaO</th>
<th>Na(_2)O</th>
<th>(\Sigma_{\text{ox}})</th>
<th>Mg#</th>
</tr>
</thead>
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<td>Pyrolite</td>
<td>45.0</td>
<td>4.45</td>
<td>0.20</td>
<td>0.04</td>
<td>0.25</td>
<td>8.05</td>
<td>0.14</td>
<td>37.8</td>
<td>3.55</td>
<td>0.36</td>
<td>99.84</td>
<td>89.3</td>
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<tr>
<td>KLB-1</td>
<td>44.5</td>
<td>3.59</td>
<td>0.16</td>
<td>0.31</td>
<td>0.25</td>
<td>8.10</td>
<td>0.12</td>
<td>39.2</td>
<td>3.44</td>
<td>0.30</td>
<td>99.97</td>
<td>89.6</td>
</tr>
</tbody>
</table>

#### Oxides in wt% and cations normalized to a total charge of 24 000 (12 000 oxygen atoms). The average compositions listed here are average values of the liquidus phases in one to seven different experimental charges. In each of the charges the liquidus phase compositions are the averages of three to 15 spot analyses, and the 1\(\sigma\) values refer to the average deviations from these mean values. The starting compositions of pyrolite \([29]\) and KLB-1 \([30]\) were prepared by combining oxides and carbonates before heating at 1200°C at an oxygen fugacity two log units below the fayalite^magnetite^quartz oxygen buffer to convert all of the ferric iron to ferrous iron. The pyrolite composition used here has about 10 times lower Cr content than that of \([29]\). In the averaging procedure used for the quenched melt and mineral analyses, only the Cr contents of the KLB-1 composition experiments are used, except for those pressures (22 and 23 GPa) where only the pyrolite composition is available. The numbers 22, 23, 24, and 24.5 denote pressure (in GPa). qm, quenched melt; fp, ferropericlase; pv, MgSi-perovskite; ga, majoritic garnet; na, not analyzed; Mg = \(100\times\text{Mg}/(\text{Mg}+\text{Fe})\), where Fe is total iron; ST = Si+Ti; CA = Al+Cr; NN = Ni+Fe+Mn+Mg+Ca+Na.
relation observed between electrical power and temperature during heating indicates that the power output is a reliable guide to the temperature. At run temperatures of 2100° to 2400°C, the thermocouple EMF often drifted slowly in spite of constant power output. Therefore, the experiments were run manually at constant power output as soon as the run temperature was reached. This resulted in far more consistent melting observations than when the thermocouple temperature was followed throughout the heating duration.

The thermal gradients within the sample capsules in comparison with the results of this melting study and previous studies (e.g. [8]) indicate that the thermal gradients at 2100–2400°C are considerably larger than those measured by two-pyroxene thermometry at 1400°C, probably on the order of 200°C/mm.

The boundary between 100% quenched melt and equilibrium crystals with interstitial melt is loosely referred to as the liquidus interface. The isotherm corresponding to this boundary, however, is not necessarily identical with the liquidus temperature, if some liquid–solid separation has occurred along the thermal gradient. Liquid–solid separation, however, is inferred to be unimportant because the quenched melt areas have fairly constant compositions similar to the bulk starting compositions (see below). The solidus cannot be observed directly in these experiments. Chemical mass balance constraints involving mineral and melt compositions, however, indicate that the appearance of Ca-perovskite is nearly coincident with the solidus in the investigated pressure range.

3. Analytical method

Recovered capsules were mounted in epoxy resin for analysis with Raman spectroscopy and electron microprobe. Two samples were also made into doubly polished thin sections for Fourier transform infrared (FTIR) spectroscopy and Mössbauer spectroscopy at the Bayerisches Geoinstitut. Electron microprobe analyses were performed in Bayreuth using a Cameca SX-50, at the Mineralogical-Geological Museum in Oslo with a Cameca Camebax Microbeam instrument and at the University of Bristol with a Jeol-8600 Superprobe. The instruments operated at an accelerating voltage of 15 kV with a beam current of 15–20 nA. The raw data were corrected by a combination of PAP and ZAF correction programs. Synthetic silicates and oxides were used as standards and the standards varied between the various microprobe locations. The minerals were mostly analyzed with a focused beam. Perovskite, which structurally decomposed under the electron beam, was analyzed both with a focused and with
a defocused beam of 5 μm diameter. No compositional difference between focused and defocused beam analyses were found, in line with the observations of Wood [26]. The damage that occurs to perovskite during analysis does not therefore seem to affect the determined composition. The quenched melt areas were analyzed partly with a raster mode of analysis covering areas of 400–700 μm² and partly by systematic and dense point transects using a defocused beam with 10 μm diameter. No systematic compositional variations were observed between the analytical data generated by the different methods and between the different laboratories.

The estimated precision and accuracy for analyses of quenched melt and different minerals, given in Table 2, are based on repeated analyses of the experimental phases and standards (10–30 analyses of each phase). Although the spatial variation of quenched melt compositions within the all-melt portions of the experimental charges are generally within the precision range, the reported melt analyses are from areas close to the liquidus. Some of the majoritic garnet crystals between the liquidus and the solidus are slightly zoned. In these cases the cores have higher Fe/Mg ratios than the rims, and they represent incompletely equilibrated subsolidus compositions formed during heating toward the run temperature. In the case of zoned crystals, the spot analyses were performed on the rims in equilibrium with the adjacent melt.

4. Results

4.1. Phase relations

The two peridotite compositions investigated have identical phase relations within the experimental resolution of this study, and there is no resolvable effect of the addition of a total of 0.55 wt% trace elements. The phase relations for these peridotite compositions at pressures of 22–24.5 GPa are shown in Fig. 1, and backscatter electron (BSE) images of representative liquidus assemblages at pressures of 23, 24 and 24.5 GPa are shown in Fig. 2. The thermal gradients across the sample capsules produce isobaric sections, covering a range of temperatures. Some of the run products cover the entire or part of liquidus to solidus range, whereas others are entirely subsolidus or supraliquidus (Table 1).

The Fe–Mg-dominated oxide phases, referred to as ferropericlase, have atomic Mg/(Mg+Fe) ratios well above 0.8, even in the subsolidus. The liquidus or near-liquidus ferropericlase has Mg/ (Mg+Fe) ratio of more than 0.9 in the entire pressure range.

Below about 23.2 GPa majoritic garnet (ga) is the liquidus phase followed by ferropericlase (fp). At higher pressures ferropericlase is the liquidus phase, followed by garnet and then perovskite (MgSi-perovskite, pv) and at 24 GPa the second crystallizing phase is perovskite followed by garnet. The crystallization of CaSi-perovskite (cpv) occurs slightly below the solidus at 22 GPa, but is nearly coincident with the solidus at higher pressures. Garnet becomes a near-solidus phase at 24.5 GPa. Recent studies of the subsolidus phase relations of peridotite compositions indicate that the gradual dissolution of garnet into perovskite with increasing pressure is completed at about 24.5 GPa at a temperature of 2100°C [26,28].

The crystallization sequences at various pressures are illustrated in Fig. 2. The BSE images demonstrate the lower thermal gradients, leading
to more interstitial melt, in the 18–8 mm assemblies (Fig. 2A,B), relative to the 10–4 mm assemblies. The crystallization sequence is inferred from the relative proximity of the different phases to the liquidus (see discussion in [8]). Three of the high-temperature run products from experiments at 24.5 GPa (Table 1, Fig. 2E) contain small amounts of ferropericlase as the only solid phase along the cold side of the sample capsules, documenting more clearly the liquidus phase at this pressure. Some of the experimental products, however, contain large and rather homogeneous quench crystals of garnet and perovskite growing from the liquidus boundary, into the liquid (Fig. 2C,D). The quenched garnet crystals in experiments 2383 and 2438 are 50–200 μm in size and occur as single and regularly zoned crystals (see analyses in Background Data Set1).

The addition of 1 wt% metallic iron to one of the 24 GPa experiments (2389) has an interesting effect on the phase relations. The resulting crystallization sequence is fp-pv-ga in contrast to the fp-ga-pv sequence observed in the other 24 GPa experiments. The stabilization of garnet relative to perovskite may be explained by the oxygen reducing effect of the addition of metallic Fe, suggesting that garnet is stabilized at more reducing conditions relative to perovskite. An ongoing experimental study carried out on a primitive mantle model composition at comparable conditions, but using diamond capsules, indicates that the crystallization of perovskite is suppressed relative to the other phases [20]. This may be explained by lower oxygen fugacity imposed by the diamond capsules (compared to Re capsules), even if the primitive mantle starting composition also has slightly higher contents of Si, Al, and Ca.

The phase relations of Fig. 1 are in broad agreement with the results of previous melting studies of KLB-1 up to 22.5 GPa [8,9], although there are minor differences. This study extends the phase relations to 24.5 GPa and documents more clearly that ferropericlase is the first liquidus phase of peridotite at pressures above 23 GPa. Although the liquidus temperature determined in our study is slightly lower than that previously determined for KLB-1 [8,9], our inferred solidus temperature is almost identical to that of previous measurements [8,27].

One of the discrepancies between previous melting studies of peridotite [8,9] and our results is the question of whether ringwoodite is present in the melting range and/or in the subsolidus assemblage at pressures of 22–23 GPa. Ringwoodite was found just above and below the solidus at about 22 GPa [9], but is absent from both the melting range and the subsolidus assemblages of all of our 22 and 23 GPa experiments. Even the 100% subsolidus experiments contain only the three-phase assemblage ga-fp-cpv at 22 GPa (experiment 2377) and the four-phase assemblage pv-ga-fp-cpv at 23 GPa (Z121).

Our results indicate that at high temperature (2000–2200°C) the perovskite-forming reaction is from majoritic garnet to perovskite as opposed to the ringwoodite to perovskite+ferropericlase reaction that occurs at lower temperatures. A recent experimental study of the subsolidus phase relations of the KLB-1 composition confirms the absence of ringwoodite near the solidus at pressures just below the perovskite-forming reaction [28]. The absence of subsolidus ringwoodite in our 22 GPa experiments is clearly not related to slow reaction rates during heating after the 20 min dwell periods at 1400°C, because the ringwoodite-out reaction boundary has a negative dp/dT slope (Fig. 1) [28].

4.2. Phase compositions

A range of mineral and melt compositions are shown in the major oxide variation diagrams in Fig. 3. Table 2 shows the average phase compositions, including normalized formula units, of minerals and melts at or slightly below the liquidus for the four different pressures. The compositions of coexisting minerals above and below the solidus are listed in the Background Data Set1.

The melt composition of each run product is homogeneous throughout the quenched melt portions of the charges and similar to the bulk composition. The melt compositions are slightly enriched in Fe and Si and depleted in Mg, relative

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1 http://www.elsevier.com/locate/epsl
to the bulk composition (Fig. 3). Generally, tie lines (not shown in the variation diagrams) between the two first crystallizing phases and the melt compositions form triangles that enclose the bulk compositions. Corresponding tie lines between the subsolidus compositions must include CaSi-perovskite, which is not included in the diagrams. The most noticeable change in mineral chemistry when going from liquidus to solidus is the strong increase in the Fe/Mg ratio. This change, observed in several other experimental melting studies (e.g. [9,11]), is seen in all of the minerals but is most pronounced in ferropericlase (Fig. 3).
The liquidus ferropericlase has a remarkably constant composition within the 22–24.5 GPa pressure range, with an Mg number \[100\times \text{Mg}/(\text{Mg+total Fe})\] of 92.3 ± 0.3 (Table 2). This rather high Mg number is slightly lower than the liquidus perovskites (94.3 ± 0.2) and garnet (94.9 ± 0.8). The Mg numbers in these minerals do not seem to vary systematically with pressure. The Mg numbers, however, decrease markedly as a function of the distance from the liquidus.

The majorite component of the liquidus garnet increases from 61.4 mol% at 22 GPa to 71.7 mol% at 23 GPa, but decreases again to 61.5% at 24 GPa, due to the appearance of coexisting perovskite. The cation sums of the garnet are within 0.2% of the expected eight cations, when normalized to a cation charge of 24 (12 oxygen atoms). The perovskite formulas, on the other hand, are characterized by small, but consistent cation excesses of 0.03–0.08 (0.4–1%) relative to the expected eight cations per formula unit of 12 O atoms. Perovskites with considerably larger cation excesses of 0.11–0.16 (1.4–2%) were observed in a similar study of a more Fe-rich composition [11].

Several studies [26,31,32] indicate that aluminous magnesium silicate perovskite has a strong affinity for ferric iron. If the entire cation excess of 0.034–0.076 in the formulas normalized to 12 oxygen atoms (liquidus perovskites at 24 and 24.5 GPa, Table 2) is ascribed to the presence of ferric iron, the charge balance of a formula normalized to 8.00 cations requires a \(\text{Fe}^{3+}/\text{Fe}_{\text{total}}\) ratio of 0.5–1. Similar magnitude \(\text{Fe}^{3+}/\text{Fe}_{\text{total}}\) ratios were measured by energy-loss near-edge structure spectrometry in aluminous perovskites equilibrated with ferropericlase and garnet in Re capsules at 24–26 GPa and 1650–1900°C (Frost and Langenhorst, submitted). The apparent cation excess may alternatively be ascribed to an aluminous- and oxygen-deficient perovskite [33]. Only 5–10% of the oxygen-deficient component \(\text{Mg}_2\text{Al}_2\text{Si}_2\text{O}_{11}\) would be required to account for the apparent cation excess.

Some of the majoritic garnet crystals have significant compositional zoning. For grains located relatively close to the liquidus the zoning involves cores with higher Fe/Mg ratios than the rims. Such Fe-rich cores represent incompletely equilibrated crystals that formed during heating towards the run temperature. Care was taken to obtain the rim composition of such grains. Similar compositional heterogeneities were not observed in perovskite and ferropericlase. The larger analytical uncertainties for garnet than for perovskite and ferropericlase (Table 2) may largely reflect these compositional heterogeneities.

The crystals of CaSi-perovskite are often too small for electron microprobe analyses. It seems that the presence of 0.45% trace element oxides in the pyrolite starting composition enhances the crystal growth. Therefore all of the analyses (Background Data Set\(^4\)) of this phase are from trace element-doped pyrolite bulk compositions. The analyzed CaSi-perovskites have major oxide totals of about 88 wt\%, implying the presence of about 12 wt% trace elements.

### 4.3. Hydrogen and ferric iron

One of the 24 GPa run products (2387) was made into a doubly polished thin section for an exploratory investigation by FTIR and Mössbauer spectroscopy. Hydroxyl was not detected at an estimated detection level of 100 ppm. Although we found no evidence for hydrogen in the crystals of the solid assemblage or quenched liquid, we cannot exclude the possibility that water in the liquid may have been quenched to a free fluid phase and thereby escaped detection.

The \(\text{Fe}^{3+}/\text{Fe}_{\text{total}}\) ratio was found to be 0.19 in the quenched melt and 0.28 in the bulk crystal aggregate. Although the iron in the starting material was reduced to ferrous iron in a gas mixing furnace, there are relatively large reservoirs of oxygen in the form of ceramic oxides in the pressure cell parts surrounding the sample capsules. Before compression these ceramic parts, as well as the sample powder itself, also contain oxygen in their pore space volumes. The ceramic parts, in particular, contain considerable pore space. It is likely that a small amount of ferrous Fe from the sample converted to metallic Fe that alloyed with the inner walls of the Re capsules during the experiments, causing some oxidation of the sample. Previous experiments in a more Fe-rich composition [11] have shown that the innermost walls of
the Re capsules can contain up to 1 wt% Fe. Due to the procedures to remove H₂O from the sample surroundings before the experiments, it is unlikely that there was any significant water present in the samples. If, however, some water was present, oxidation of the sample could conceivably occur by dissociation of H₂O in combination with diffusive H loss.

It is interesting to note that the proportion of ferrous Fe is higher in the bulk crystal aggregate than in the quenched melt. The reason for such a preferential partitioning is most likely that Al-bearing perovskite has greater affinity for Fe³⁺ than the other phases, including the quenched melt phase (see the discussion above).

**5. Mineral–melt partitioning**

5.1. Major elements

Fig. 4 summarizes the mineral–melt partitioning of major and minor elements from our study and includes some of the data from Herzberg and Zhang [9]. There is a good agreement between our determined partitioning ratios and those reported by [9]. In Fig. 4 the mineral–melt partitioning of major elements is shown as exchange distribution ratios, labelled $K_D$, involving the Si/Mg, Fe/Mg and Ca/Al ratios. These $K_D$ values [=$(i/j)_{\text{mineral}}/(i/j)_{\text{melt}}$] do not correspond to chemical equilibrium constants for simple ionic exchange reactions between solid and liquid, and are therefore not strictly equivalent to true exchange distribution coefficients [34]. This applies even to the Fe/Mg ratio, because of the complex Fe partitioning in perovskite (e.g. [32]). Hanson and Langmuir [35] classified these elements as essential structural constituents (ESC: Si, Al, Ca) and intermediate elements (Mg, Fe⁵⁺), respectively. The ratios of true ESC elements are constant for a given mineral, and a variation of exchange distribution coefficients for ESCs will therefore reflect the variation in melt composition. The increasing solubility of the majorite (Mg,Fe)SiO₃ component in garnet with increasing pressure, however, implies that Si, Al and Ca cannot be considered true ESC elements in this phase.

At lower pressures the mineral–melt $K_D$(Fe/Mg) is largely independent of composition and temperature, because of the geochemical similarity between these two elements and because the compositional effects on the $D_i$ (Nernst distribution coefficient) for Fe and Mg cancel out [9]. Such a simple Fe–Mg relationship does not exist for perovskite, however, due to the preferential

![Fig. 4. Mineral–melt partitioning ratios for major and minor elements. Symbol shapes as in Fig. 3, small symbols from Herzberg and Zhang [9] and large symbols from this investigation. The $K_D$ ratios are not true partitioning coefficients linked to equilibrium constants of simple cation mineral–melt exchange reactions (further discussion in the text). The $D_{Cr}$ values derived from our data set are based on only the experiments with KLB-1 starting material. $K^2_{ij} = (i/j)_{\text{mineral}}/(i/j)_{\text{melt}}$; $D_i = (i)_{\text{mineral}}/(i)_{\text{melt}}$. The propagated 2σ error bars, shown to the right of each of the panels, are calculated from the average analytical errors in Table 2.](image)
incorporation of significant amounts of ferric iron in a coupled substitution with Al. In Fig. 4 the Fe/Mg ratio involves the sum of ferric and ferrous Fe, even if Fe$^{3+}$ is partitioned selectively into the bulk mineral assemblage relative to the liquid, and into the perovskite relative to garnet and ferropericlase.

The $K_D$(Si/Mg) for garnet remains almost constant at 1.3 in the 20–24 GPa range, whereas the corresponding value for perovskite is slightly lower (1.2) and shows a decreasing trend from 24 to 24.5 GPa. The $K_D$(Fe/Mg) remains well below unity for all of the minerals. With increasing pressure from 20 to 25 GPa, an increase from below 0.4 towards 0.5 for garnet and perovskite is opposed by a decrease from 0.7 to 0.6 for ferropericlase. The increase in the Fe/Mg ratio of the melt relative to the bulk solid residue is accompanied by an even stronger increase in the Ca/Al ratio of the melt. The $K_D$(Ca/Al) is well below 0.4 for all of the minerals. With increasing pressure, the $K_D$(Ca/Al) increases for garnet and decreases for perovskite. The major element partitioning coefficients derived for the investigated peridotite compositions overlap with corresponding coefficients for a composition with much higher Fe/Mg ratio [11], indicating that the $K_D$ values are rather insensitive to moderate variations in bulk composition.

5.2. Minor elements

The mineral–melt partitioning of the minor elements, Ti, Cr, Ni, and Na, is shown by the simple Nernst distribution coefficients in Fig. 4. Ti is compatible in perovskite ($D_{\text{Ti}} = 1.3–1.4$) and incompatible in garnet and ferropericlase ($D_{\text{Ti}} < 0.1$). The $D_{\text{Ti}}$ value for garnet increases from 0.2 to 0.6 with increasing pressure. The behavior of chromium is inverse to that of Ti, with $D_{\text{Cr}}$(garnet) decreasing from 1.5 to 1.1 and $D_{\text{Cr}}$(ferropericlase) decreasing from 1.8 to 1.4 with increasing pressure. Cr is weakly incompatible in perovskite. Nickel is strongly compatible in ferropericlase and incompatible in garnet and perovskite, with partitioning coefficients of 4.0–4.3 and 0.2–0.4, respectively. There is no discernible variation in the Ni partitioning as a function of pressure. Sodium is incompatible in all of the solid phases. The compatibility in both garnet and ferropericlase, however, seems to increase as a function of pressure, and the $D_Na$ value for ferropericlase reaches 0.6 at 24.5 GPa.

It is interesting to note that the weak pressure-dependent trends for garnet–melt partitioning of Ti, Cr and Na observed in Fig. 4 are largely in accordance with the garnet–melt partitioning at 15 GPa [12]. Various other studies of garnet–melt and perovskite–melt partitioning, however, show somewhat different behavior for some of these elements. Whereas Drake et al. [36] and McFarlane et al. [43] report $D_{\text{Cr}}$(garnet) values of 1.0 to 1.5, which are broadly in agreement with our results, Ohtani et al. [37] derived a higher value of 2.3. The $D_{\text{Cr}}$(perovskite) values and $D_{\text{Cr}}$(perovskite) values of [13,14,36,38–41] have wide ranges of 0.6–3.0 and 0.4–6.3, respectively. The values of McFarlane et al. [13], however, are in overall agreement with the present results. There are considerable differences in major element bulk composition between the calcic ultrabasic system of [14,40] and the peridotitic composition of the present study. The Ni partitioning perovskite–melt and ferropericlase–melt, however, does not show large variations between the various studies, and the ferropericlase–melt partitioning of Ti and Cr is generally also consistent between the studies.

The recent development of improved microanalytical techniques for trace element analyses (laser ablation ICPMS and SIMS) makes it possible to acquire a large range of elemental mineral–melt partitioning data. The design and performance of stable equilibrium experiments at pressures above 20 GPa now represent the main challenge in generating high-quality partitioning data for minor and trace elements between majoritic garnet, ferropericlase, perovskite and ultramafic silicate melts.

6. Implications for early mantle differentiation

The liquidus phase relations of common peridotite compositions derived in this study demonstrate clearly that ferropericlase is on the perido-
tite liquidus above about 23.2 GPa. This was not recognized in some of the early melting studies of peridotitic and chondritic composition in this pressure range (e.g. [7,42]). Later, studies (e.g. [8,10,11,43]), however, have observed ferropericlase on or near the liquidus in peridotitic and chondritic compositions at pressures corresponding to the transition zone and the lower mantle. In this study, three of the high-temperature run products from experiments at 24 GPa contain small amounts of ferropericlase as the only solid phase along the cold end of the sample capsules, demonstrating clearly that ferropericlase is the first liquidus phase. Although ferropericlase is the liquidus phase in the 23–25 GPa range, ongoing melting studies, using multianvil configurations with sintered diamond anvils, indicate that perovskite becomes the first liquidus phase above 31 GPa [44]. At 33 GPa Ca-perovskite also crystallizes closer to the liquidus and almost simultaneously with ferropericlase.

Another important observation from this investigation is the absence of ringwoodite in the high-temperature part of the subsolidus region at pressures just below the perovskite-forming reaction boundary. This implies that the high-temperature perovskite-in transition has a positive \( \frac{dp}{dT} \) slope [28] and that hot plumes ascending from the lower mantle will be accelerated rather than delayed at the bottom of the mantle transition zone.

An implication of the early crystallization of ferropericlase from melts in the lower part of the transition zone and the uppermost part of the lower mantle is that the proposed mechanism of producing a lower mantle with an elevated Si/Mg ratio relative to the upper mantle and chondritic compositions by fractionation of perovskite [45] may not be a viable process. Because the observed \( K_D(Si/Mg) \) for perovskite is close to unity (1.2), the addition of even small amounts of ferropericlase to the crystallizing assemblage would decrease rather than increase the Si/Mg ratio in bulk solid material. At depths exceeding 850 km, however, the possibility of perovskite-dominated fractionation [44] may facilitate the production of residues with elevated Si/Mg ratios relative to a bulk silicate Earth composition. The additional possibility of near-liquidus coprecipitation of Mg- and Ca-perovskites may further enhance the residual Si/Mg ratio, as well as the REE and HFSE concentrations in the residue or fractionated assemblage.

The density relations between coexisting perovskite, ferropericlase and melt at 500–800 km depth in the Earth [11,45,46] indicate that partial melts will be buoyant relative to these minerals, and that fractionation of ferropericlase and perovskite from rising melt fractions, either in a solidifying magma ocean or from subsequent hot plumes, can be expected to occur. Strong convective motions during magma ocean crystallization, however, may largely have erased the memory of such a fractionation [2].

The partitioning relations shown in Fig. 4 indicate that fractional crystallization or partial melting in the transition zone and the upper part of the lower mantle would increase the Fe/Mg and Ca/Al ratios and the Na concentration of the melt. Whereas the \( K_D(Fe/Mg) \) for ferropericlase–melt partitioning is always lower than 0.7 in our measurements and is observed to decrease with pressure towards lower mantle conditions, previous studies [47,48] have found above-unity values for this partitioning. Such discrepancies may result from the measurement of subliquidus ferropericlase compositions in extreme thermal gradients [11].

The mineral–melt partitioning of Ti and other high field strength elements is strongly dependent on whether perovskite is part of the residual or fractionating assemblage. The high field strength elements are compatible in perovskite, but incompatible in garnet and ferropericlase. Cr partitioning is generally opposite to that of Ti, although the \( D_Cr \) for all of these phases cluster closer to unity than the \( D_{Ti} \).

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