The role of sulphur during partial melting of eclogite in the cratonic mantle: constraints from experiments and mantle xenoliths

Sara Burness¹, Katie A. Smart¹, Gary Stevens², Sebastian Tappe³

¹ School of Geosciences, University of the Witwatersrand, Johannesburg, South Africa; sara.burness1@gmail.com, katie.smart2@wits.ac.za
² Centre for Crustal Petrology, Stellenbosch University, Stellenbosch, South Africa, gs@sun.ac.za
³ University of Johannesburg, Auckland Park, South Africa, sebastian@uj.ac.za

Introduction

Sulphide- and carbon-bearing melts play an important role as metasomatic agents within the cratonic mantle lithosphere, and may be linked to processes including diamond formation (Griffin et al., 2003), metal transport (Arndt et al., 2005) and the deep carbon cycle (Tappe et al., 2017). While the influence of C-bearing fluids on the lithospheric mantle is under continuous investigation (e.g. Dasgupta et al., 2004), the activity of S-bearing fluids and their role in diamond formation is less well constrained (Mungall and Brenan 2014). Sulphide minerals are key to the mantle S cycle. They control the platinum group element (PGE), siderophile and chalcophile element budgets, and their behaviour during partial melting controls the movement of these economically important metals. Here we present a case study of eclogite metasomatism by S+C-bearing fluids, by comparing new experimental work with geochemical investigations of sulphide-bearing eclogite xenoliths from several kimberlite occurrences on the Kaapvaal craton, South Africa. The focus of this study is to delineate the activity of S on the melting behaviour of eclogite and the relationship between sulphide-melt formation relative to carbonate+silicate melt formation. Our findings add to the knowledge of volatile-bearing solidii in the upper mantle, the partitioning nature of various major and trace elements between the silicate residuum, possible sulphide residuum, and miscible or immiscible silicate+carbonate melts and S-melts. A complementary investigation of sulphide-bearing eclogite xenoliths may serve as a benchmark for the processes of volatile-bearing metasomatism studied here.

Melting of C+S-bearing MORB at upper mantle conditions

In order to investigate the influence of C-O-H-S bearing fluids on mantle processes, we present results from partial melting experiments that constrain the near-solidus phase relations of S- and C-bearing eclogite from 2.0 to 3.5 GPa and 1050 to 1300°C. A synthetic MORB-like composition created from reagent grade silicate and carbonate powders was fired into a glass at 1400°C before being doped with 5wt.% synthetic pentlandite (Fe₀.₉₃Ni₀.₁₇Cu₀.₀₃S₁.₁₀; Metal/Sulphur = 1.12; Ni/(Fe+Ni) = 0.15) and, to foster melt detection, various proportions of CO₂ (0.5 and 15wt.%) in the form of CaCO₃. The starting compositions were placed in graphite-lined PdAg capsules and experiments were conducted using non-end loaded piston cylinder apparatus with a run time of 24 hours. The silicate experimental products consist of euhedral to subhedral garnet and omphacitic cpx, +coesite, +zircon and +anorthite (Fig. 1), typically ≤15µm in size. Both sulphide liquid (e.g., Fe₁.₀Ni₀.₁Cu₀.₀₆S₁.₁₂; Metal/Sulfur = 0.93; Ni/(Fe+Ni) = 0.09) and MSS (e.g., Fe₀.₇₅Ni₀.₂₂Cu₀.₀₅S₁.₂₆; Metal/Sulphur = 0.84; Ni/(Fe+Ni) = 0.22) are precipitated during the partial melting of eclogite. Phase relationships (Fig. 1) show distinct rounded globules of sulphide melt (L₃) which are texturally immiscible to both silica and silica+FeS dominated melts. MSS form discrete sub-angular grains that are usually ≥15µm in size. The location of the sulphide solidus is plotted in Fig. 2 along with carbonated eclogite solidi (1 and 2), a CO₂-rich vapour is inferred as the main medium of carbon at low pressure in experiments with 5wt.% CO₂.
Figure 1: Select BSE images of the experimental run products. Cpx, clinopyroxene; Omp, omphacite; Diop, diopside; Grt, garnet; Zrn, zircon; An, anorthite; CC-Dol, calcio-dolomitic solid solution; MSS, massive sulphide solid solution; Melts, L_{Si} = silica-rich liquid, L_{Si,C} = silica-carbonate liquid, L_S = sulphide liquid.

Figure 2: Experimental P-T phase diagram for S+C-bearing MORB. Phase abbreviations as in Fig. 1. Black fields indicate the presence of a mineral phase, red fields a quenched liquid phase (melt), blue fields ≤ 2vol. % of a liquid phase plus voids inferred to represent CO_2-gas pockets. Sulphide solidus and liquidus curves are plotted using thick solid lines. 1. carbonated eclogite solidus (15wt.% CO_2) and 2. carbonated eclogite solidus (5wt.% CO_2). EC1 (15wt.% CO_2; Yaxley and Brey, 2003), SLEC1 (5wt.% CO_2; Dasgupta et al. 2004), CO_2-free solidus (CO_2-free eclogite, Litasov and Ohtani 2010).

Figure 3: Reconstructed bulk eclogite xenolith sulphides and experimentally produced MSS from this study plotted in a Fe-S-(Ni+Co) ternary diagram. Grey shaded area represents sulphides from eclogite xenoliths and E-type diamond sulphide from the Kaapvaal Craton. The grey circle encapsulates sulphides which have been affected by oxidation.

Textural and compositional characteristics of sulphides in mantle eclogite xenoliths

Nineteen eclogite xenoliths from the Jagersfontein, Roberts Victor, Kamfersdam, Swartruggens and Premier kimberlites were selected for study based on the identification of sulphides. Sulphides form discrete polyphase assemblages (e.g. Po+Pn+Cp) that occur as inclusions in clinopyroxene along with interstitial grains. The reconstructed sulphides are similar in composition to some sulphides present in mantle eclogites, eclogite-type diamond inclusions as well as the MSS produced in our experiments (Fig. 3). The natural sulphides (MSS) can contain up to hundreds of ppm Mo, Zn, Ag, and Pb, <100 ppm of Ag and V, whereas Pd, Ru, Rh, Pt, and Au concentrations are <10ppm. Os and Ir are present from bdl to < 0.3ppm and <0.6ppm, respectively. Zn, Mn and Co are the most abundant of the sulphide trace elements (e.g. Zn ranges in concentration from ~1.8 ppm to ~1 wt%). Most eclogite xenoliths studied here have oceanic crustal protoliths, based on normalized REE patterns, Eu anomalies, and oxygen isotope compositions (δ^{18}O values from 3.3 to 10.4‰).
Links to mantle metasomatism in the presence of C+S- bearing fluids/melts

Our study shows that S melts+MSS are the first to appear upon melting of a S+C-bearing high-pressure MORB (Fig. 2). There is a small temperature difference between the S and S+C solidii at the pressures investigated, but this temperature difference may not easily be distinguished during incipient melting. The experimental S-melts and solid MSS are enriched in siderophile and chalcophile elements and likely PGEs (Bockrath et al., 2004), show a striking similarity to sulphide inclusions in E-type diamonds (Fig. 3: Deines and Harris, 1995) and texturally and compositionally resemble the sulphides in our studied eclogite xenoliths. The experimental S-melts and MSS also appear to form high-angle contacts with silicates and Si+C melts, or discrete globules within highly polymerised silicate melt (Fig. 1), such that the immiscible nature of our experimental MSS supports the ‘nugget effect’ of sulphides observed in mantle eclogite nodules. In summary, our study suggests that S-melts alone may not be effective metasomatic agents in the cratonic mantle. However, since S+C melts are likely produced together during mantle melting, carbonatic melts with appreciable S may instead be effective ways to concentrate and mobilize PGE minerals within the CLM, and due to the overlap of sulphide DIs and our experiments, S+C melts may be effective diamond-growth media.

References

Arndt N, Lesher CM, Czamanske GK (2005) Mantle-derived magmas and magmatic Ni-Cu-(PGE) deposits: 5-24
Yaxley GM, Brey GP (2004) Phase relations of carbonate-bearing eclogite assemblages from 2.5 to 5.5 GPa: implications for petrogenesis of carbonatites. Contributions to Mineralogy and Petrology 146(5): 606-619