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Continuous eclogite melting and variable refertilisation in upwelling heterogeneous mantle

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Large-scale tectonic processes introduce a range of crustal lithologies into the Earth's mantle. These lithologies have been implicated as sources of compositional heterogeneity in mantle-derived magmas. The model being explored here assumes the presence of widely dispersed fragments of residual eclogite (derived from recycled oceanic crust), stretched and stirred by convection in the mantle. Here we show with an experimental study that these residual eclogites continuously melt during upwelling of such heterogeneous mantle and we characterize the melting reactions and compositional changes in the residue minerals. The chemical exchange between these partial melts and more refractory peridotite leads to a variably metasomatised mantle. Re-melting of these metasomatised peridotite lithologies at given pressures and temperatures results in diverse melt compositions, which may contribute to the observed heterogeneity of oceanic basalt suites. We also show that heterogeneous upwelling mantle is subject to diverse local freezing, hybridization and carbonate-carbon-silicate redox reactions along a mantle adiabat.

he plate tectonic paradigm suggests that oceanic crust and underlying lithospheric mantle are recycled into the mantle as cool, subducted slabs at convergent margins. During subduction of oceanic lithosphere, basaltic crust reacts to form quartz/coesite eclogite at high pressure and low temperature conditions under which partial melting is avoided or limited to low temperature melting at the wet solidus (that is the temperature at which a silicate melt first appears in a rock in the presence of an aqueous fluid). The breakdown of hydrous phases such as lawsonite and phengite results in the liberation of an aqueous fluid during progressive metamorphism but traces of water can be retained in eclogitic garnet and clinopyroxene that are nominally anhydrous minerals (NAMs)¹⁻⁵. Subducting slabs often descend to the Transition Zone, and in some cases, into the lower mantle^{6,7}. As such slabs are dominated by residual to refractory lithospheric peridotite (depleted lherzolite to harzburgite), their negative buoyancy is reversed if their temperatures approach that of surrounding fertile peridotitic mantle ('fertile' MORB-source mantle or 'modern, well-mixed mantle'). Heterogeneity in the mantle persists when subducted slabs dominated by depleted peridotite with discrete bodies of quartz/coesite eclogite (former oceanic crust) are entrained and mixed into convecting mantle. Later sampling of such mantle domains during partial melting is regarded as essential to the formation of some mantle-derived primitive magmas⁸⁻¹³. The model being explored here postulates that subducted slabs heated to approach the intraplate geotherm will undergo localised melting within slab heterogeneities or at slab/fertile mantle interfaces. As temperatures equilibrate, buoyant adiabatic upwelling of heterogeneous peridotite mantle containing discrete bodies of coesite eclogite induces local melting and possible metasomatism of surrounding ambient mantle.

Spandler et al.¹² presented the melting phase relations of a model composition (GA2) representative of fertile oceanic crust over a decompression path from ~ 160 km (5 GPa). A composition (Res2) of the residue after extraction from GA2² of $\sim 15\%$ potassic dacitic-rhyodacitic melt at 5 GPa forms the basis of our experimental study. It is a 'dry' (nominally anhydrous) coesite eclogite (Supplementary Fig. (SFig.) 1; Supplementary Table (STable) 1). In our experiments we investigated further temperature rise at 5 GPa with consequent melting of Res2. We also crystallised Res2 under conditions simulating further decompression melting from ~ 160 to

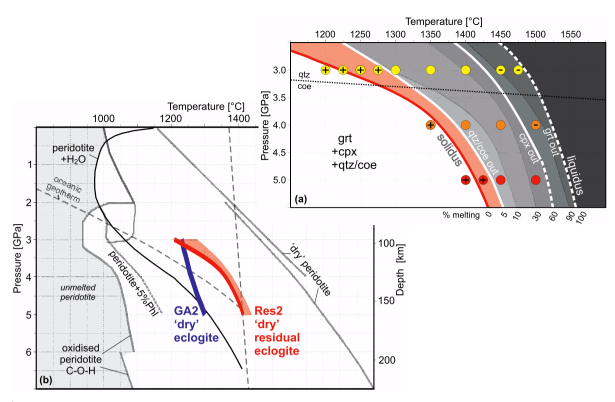


Figure 1 (a) Pressure-Temperature-grid of experimentally-determined melting and phase relations of Res2. (b) Comparison of Res2 solidus surface and its (incipient) melting regime with solidii of other experimentally determined mantle lithologies. (a) See explanations within text. Yellow circles – Res2 at 3 GPa, orange circles – Res2 at 4 GPa, red circles – Res2 at 5 GPa. The symbols '+' and '-' indicate the presence of quartz/coesite and absence of clinopyroxene, respectively, as residual phases. Solid and dashed white lines are determined and inferred phase boundaries, respectively. The quartz-coesite transition is taken from Bose & Ganguly⁴⁴; qtz, quartz; coe, coesite; grt, garnet; cpx, clinopyroxene. (b) The Res2 solidus lies between the 'dry' lherzolite solidus^{29,45}, and GA2 eclogite¹² and volatile-bearing lherzolite (i.e. peridotite + H₂O ± CO₂) solidus^{26,29,30–32}. The light red shaded area adjacent to Res2 solidus is the incipient melting regime with garnet + clinopyroxene + quartz/coesite residue. Predicted mantle T_p's range from 1280–1600°C^{46–48} but for illustrative purposes, we chose a mantle adiabat of T_p ≈ 1360°C so that the solidus of Res2 intersects the adiabat at 5 GPa at ~1410°C, initiating melting. Using other adiabats will simply shift the pressure of the onset of melting of various lithologies by ~1 GPa per 100°C¹². Although such changes would not alter the principles of the scenarios¹², they would substantially influence the composition of melts and solid residues of mantle lithologies at the onset of melting, and thus, would also influence any associated mantle metasomatism.

 \sim 90 km depth (5 to 3 GPa) to examine melts and residues along an upwelling path (STable 2). Our guiding concept is that these local eclogite-derived melts migrate into, and react with, enclosing peridotite/harzburgite.

Results of Res2 phase and melting relations at 3-5 GPa, 1200-1500°C are presented in Figs. 1–3, and in the supplementary items (Supplementary Notes (SNotes), SFigs. 1-11, and STables 1-4). The initial aim of this research was to perform experiments under anhydrous conditions. However, analyses of glasses from quenched melts in several runs, especially in those with low melt fractions, often returned very low totals (down to 92.3 wt.%; STable 3), indicative of the possible presence of a volatile component in the quenched liquid. Therefore, experiment Res2_19 containing 99% melt was subject to quantitative Fourier-transform infrared spectroscopy (FTIR) analysis, and ~0.27 wt% H2O was detected (SFig. 2; STable 4). Hence, the Res2 starting composition is expected to contain ~0.26 wt.% of H₂O. This water content is of the order that may be contained in nominally anhydrous ('dry') residual eclogite², but its presence demonstrates that hydrogen could not be entirely eliminated from the experiments and must be considered during interpretation of the results, as the presence of water may lower solidus temperatures relative to truly anhydrous runs. Subsolidus phases for Res2 are garnet, omphacitic clinopyroxene and quartz/coesite (Fig. 1a; SFigs. 3 & 4; STables 2 & 3; SNotes). The Res2 solidus is located at 1210 \pm 15° C at 3 GPa, $1375 \pm 25^{\circ}$ C at 4 GPa, and $1410 \pm 15^{\circ}$ C at 5 GPa (Fig. 1; STable 2). At very low degree (incipient) melting, liquids appeared mainly at triple grain boundaries as tiny (\leq 5 µm), partly connected pools of quenched glass in assemblages where quartz/coesite is present (SFig. 3; STable 2). As in previous studies¹²⁻¹⁵, quartz (\leq 3.0% at 3 GPa) or coesite (<1% at 5 GPa) is a residual phase in association with a eutectic-like incipient (\leq 5%), dacitic and sodic melt near the solidus over a temperature interval of up to 100°C at 3 GPa, but <50°C at 5 GPa (Fig. 1; SFigs. 3–6; STable 2). As quartz/ coesite is melted out, the melt fraction increases, the Mg* [100 × Mg/ (Mg + Σ Fe)] of garnet and clinopyroxene increases, and the melt compositions move along garnet-clinopyroxene cotectics, becoming progressively less siliceous and less sodic (Fig. 1; SFigs. 3–9; STables 2 & 3). The clinopyroxene/garnet ratio of the residues decreases with increasing melt fraction until clinopyroxene is exhausted, leaving garnet as the sole liquidus phase (Fig. 1a; SFig. 4; STable 2).

Residual garnet and clinopyroxene solid solutions are similar to those from other experimental studies of quartz/coesite-eclogite¹²⁻¹⁶ (SFigs. 6–9; SNotes). Significantly, clinopyroxene contains a substantial Ca-eskolaite component as a consequence of reaction between excess SiO₂ (quartz/coesite, or SiO₂-rich melt) and Ca-Tschermaks solid solution in clinopyroxene¹²⁻¹⁷ (Fig. 2; SFig. 9; STable 4; reaction 1).

$$2Ca_{0.5}[\blacksquare]_{0.5}AlSi_2O_6 = CaAlAlSiO_6 + 3SiO_2$$
(1)

Ca - eskolaite = Ca - tschermaks + melt/quartz/coesite,

where "■" represents a vacancy on the M2-site.

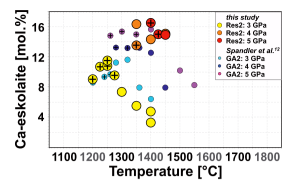


Figure 2 | Ca-eskolaite component in experimentally crystallised clinopyroxene from Res2 and GA2 as a function of pressure and temperature. The symbol '+' indicates the presence of quartz/coesite as residual phase. At a given pressure, both below and above quartz/coesite out, Ca-eskolaite first increases and then decreases with increasing temperature. Although melting residues of Res2 lose siliceous melt fractions and become increasingly depleted in Ca-eskolaite, they still retain their SiO₂–saturated character (as will *their* resulting melt fractions), and are not exhausted in Ca-eskolaite during continuous melt extraction on adiabatic ascent (this Fig., SFig. 6; STable 3). Thus, in contrast to previous predictions²⁰, melting residues of initial coesite/quartz-bearing or bimineralic eclogite with Ca-Eskolaite component in clinopyroxene present never become SiO₂-deficient (in terms of high pressure normative components) along adiabatic paths, and remain on the SiO₂-excess side of the thermal divide in SFig. 6.

The high temperature decrease in Ca-eskolaite (Fig. 2) corresponds to a decrease of silica activity towards higher temperature as soon as quartz/coesite is melted out, driving reaction 1 to the right. However, as long as the activity of silica is unity, i.e. as long as quartz/ coesite is present, this reaction will move to the left.

The increased Ca-eskolaite component in clinopyroxene at greater mantle depths causes melting to occur over a much smaller temperature interval in association with coesite at 5 GPa than with quartz at 3 GPa (Figs. 1 & 2; SFig. 4). In addition, towards higher pressures the grossular component in garnet and the jadeite component in clinopyroxene are stabilized, whereas the Ca-Tschermaks (Ca-Al-rich) component and the Al-buffonite (Ti-rich) component of clinopyroxene are destabilized (SFigs. 7–9).

During upwelling of heterogeneous mantle along an adiabatic path, residual eclogite Res2 starts to melt at much higher pressure than surrounding 'dry' peridotite, as has been also suggested for other nominally anhydrous eclogite compositions^{8,10,12-14,18,19} (Fig. 1b). Taking the potential temperature (T_p) of the upwelling mantle to be 1360°C, incipient melting of Res2 begins at 5 GPa (~160 km depths) at 1410 \pm 15°C (Fig. 1b), consistent within experimental error with derivation of the Res2 composition by extraction of ~15% melt from GA2 at 1380°C at 5 GPa. The relative slopes (dT/dP) of the adiabat and the Res2 solidus (dT/dP = \sim 30- 40° C/GPa from 5 to 4 GPa, and dT/dP = $\sim 160-170^{\circ}$ C/GPa from 4 to 3 GPa) ensure that adiabatic upwelling of Res2 leads to a selffluxing, continuous melting process (Fig. 1b). This is because adiabatic ascent of Res2 from ~160 to ~120 km depths (5 to 4 GPa) results in the continuous breakdown of Ca-eskolaite solid solution in clinopyroxene, causing exsolution of SiO₂ as both coesite and SiO₂rich melt fractions, and relative increase of Ca-tschermakite solid solution in residual clinopyroxene (reaction 1; Figs. 1 & 2; SFigs. 8 & 9; STable 3). Consequently the solidus and melt compositions remain at the garnet + clinopyroxene + coesite eutectic-like field with continuous production of highly siliceous, incipient melt fractions (reaction 1; Figs. 1 & 2; SFigs. 5 & 6).

This distinctive style of self-fluxed, eutectic-like melting rapidly increases during further adiabatic ascent of Res2 from \sim 120 to

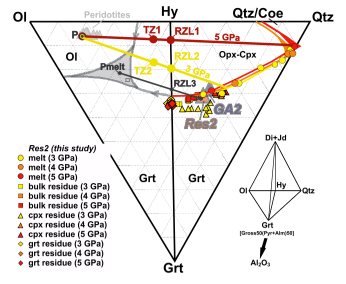


Figure 3 | Hybridisation reactions between eclogitic Res2 partial melts and peridotite in the pseudo-ternary Ol-Qtz-Grt diagram projected from (Di + Jd). Dark red and dark yellow lines illustrate normative compositions of "reaction-zone lithologies" (RZL) formed by reaction between peridotite and incipient, highly siliceous (dacitic) and higher degree, less siliceous (andesitic to basaltic) Res2 melts at 5 GPa (RZL1) and 3 GPa (RZL2), respectively. This leads to a higher normative garnet/ orthopyroxene ratio and more Cr-rich clinopyroxene in RZL2, as 3 GPa Res2 melts are less siliceous and more sodic than 5 GPa Res2 melts (see text; SFigs. 5-6, 10; reactions 1-2; reactions 1-3 in SNotes). The compositional variations within the orthopyroxene RZLs formed at different pressures along an adiabat lie on the Hy-Grt thermal divide and lead to corresponding compositional variations in the transition zone layers (TZ1 and TZ2, i.e. the olivine orthopyroxenite/garnet websterite boundaries that mark the silica-undersaturated reaction zone boundaries between peridotite [harzburgite/lherzolite] and the orthopyroxene-rich RZLs). The resulting melt-depleted residual eclogite zones vary considerably in composition over the pressure and temperature range of interest. The dark grey line expresses the hybridisation reaction of incipient melts of peridotite (+H2O) with solid 'dry' residual eclogite Res2. This join Grt-Hy represents a thermal divide of garnet-pyroxene¹³. Ol, olivine; Qtz, quartz; Coe, coesite; Grt, garnet; Hy, hypersthene; Di, diospide; Jd, jadeite; ss, solid solution; P, peridotite; M, eclogitic melt; RZL, orthopyroxene-rich reaction zone layer; TZ, transition zone layer (i.e. the contact between peridotite and orthopyroxene-rich RZL).

~90 km (4 to 3 GPa), due to the sharp increase in exsolution of coesite/quartz from Ca-eskolaite-bearing clinopyroxene, which causes steepening of the Res2 solidus slope, enhancing melt production (Figs. 1 & 2; SFig. 4). Melting at 3 GPa therefore occurs at lower temperature and yields much higher fractions of less siliceous melt than at deeper mantle levels (5–4 GPa) along the adiabatic path (~5–10% at 4 GPa to ~35% at 3 GPa) (Fig. 1; SFigs. 4–6).

The destabilisation of Ca-Eskolaite, jadeite, and grossular but stabilisation of Ca-Tschermaks and Al-buffonite during mantle upwelling causes decreased partitioning of Na and Al but increased partitioning of Ca and Ti into clinopyroxene and decreased partitioning of Ca into garnet relative to melt (SFigs. 7–10; STable 3). Higher Res2 melt fractions (>15–20%) formed at 3 GPa are therefore more sodic and aluminous, but less siliceous and titanifereous than incipient Res2 melts formed at 4–5 GPa along the chosen adiabat (Fig. 1; SFigs. 5 & 6). Consequently, as the residues of Res2 melting at 3 GPa lose ~15 to \leq 35% melt, they become bimineralic eclogite (residual garnet and clinopyroxene only) with relatively high Mg[#]. Clinopyroxene is characterised by high Ca-tschermaks and diopside, and relatively low Ca-eskolaite and jadeite components,

relative to clinopyroxene from coesite-eclogite residues of Res2 melting at 5–4 GPa with \leq 5% melt loss during adiabatic upwelling (Fig. 2; SFigs. 8–9; STable 3). Residual garnet at 3 GPa has a higher pyrope and lower almandine and grossular components with respect to residual garnet at 5 GPa (SFig. 7).

During adiabatic upwelling of heterogeneous mantle from 5 to 3 GPa, ambient peridotitic mantle will be metasomatised by (highly) siliceous melt fractions continuously extracted from Res2 eclogite. These melt fractions will react with peridotite mineral phases (mainly olivine but also pyroxene and garnet), and thus, freeze into the ambient mantle, producing an orthopyroxene-rich reaction zone layer (RZL) with garnet and minor clinopyroxene lying on the thermal divide¹³ (Fig. 3; reaction 2; reactions 1–3 in SNotes). Mineral compositions and proportions of the newly formed RZL's will reflect some admixtures of peridotite and varying siliceous and sodic melt compositions produced by continuous melting of Res2 during adiabatic ascent (Fig. 3). This process therefore adds eclogite derived incompatible elements back into an ambient (depleted) peridotite mantle. As the extracted melt compositions vary, metasomatic effects will also vary along the adiabatic path (Figs. 1 & 3).

The behaviour of Na₂O and Cr₂O₃ are potentially useful in discerning the effects of melting and reaction in heterogeneous eclogite/ peridotite lithologies. Cr₂O₃ is refractory with respect to eclogitic melts. The Cr/Al of oceanic crust is relatively low ($<0.01^{12,17,20}$). Residual garnet and clinopyroxene remain low in Cr₂O₃ and Cr/Al even after extraction of high melt fractions from residual eclogite^{12,17,20}. Cr/Al in fertile lherzolite is much higher and increases in residues from partial melting²¹, so highly refractory harzburgite residues have the highest Cr/Al ratios and Mg#s²¹. In contrast, Na₂O and Al₂O₃ are mobile in eclogitic partial melts (GA2 and Res2) but are controlled by partitioning relationships with clinopyroxene in residual eclogite (SFigs. 5 & 10; STable 3). High % Res2 melt fractions at 3 GPa are more enriched in Na₂O than 5 GPa incipient melts (SFig. 5) and are most effective in leaching Cr³⁺ and Fe³⁺ from garnet of adjacent peridotite (or reacting with chromite of adjacent harzburgite or dunite), relative to Res2 low-degree melts formed at 5 GPa. Therefore, reaction between eclogitic melts and peridotitic mantle at relatively low pressure (e.g. 3 GPa) is more effective than at high pressure (e.g. 5 GPa) at forming Na-Cr-rich (and Na-Fe³⁺-rich) pyroxene such as kosmochlor (and acmite) in the RZL according to reaction 2 (and reaction 3 in SNotes).

$$\begin{split} (Ca,Mg)_{3}(Cr,Fe^{3+})_{2}Si_{3}O_{12}+2NaAlSi_{2}O_{6} = \\ 2Na(Cr,Fe^{3+})Si_{2}O_{6}+(Ca,Mg)_{3}Al_{2}Si_{3}O_{12} \end{split} \tag{2}$$

 $Cr - Fe^{3+}$ garnet (peridotite) + jadeite (eclogitic melt)

=kosmochlor, acmite (RZL) + grossular (RZL).

Reports of Na + Cr-rich pyroxene at the contacts between refractory eclogite layers in peridotite in peridotite-pyroxenite massifs²², as well as frequently observed 'eclogitic' garnets containing little or no Cr and Na included in some diamonds²³, may form via metasomatic processes rather than extensive melt depletion of peridotite, or garnets crystallised from mafic (i.e. eclogitic) lithologies, respectively^{22,23} (see also below).

Assuming a Res2 bulk H₂O content of ~0.26 wt.%, Res2 melt fractions will release considerable H₂O during adiabatic upwelling (a maximum of 4 wt.% H₂O in melt at 5 GPa for a 5% melt; ~0.5 wt.% H₂O in melt at 3 GPa for a 40% melt). This results mainly from the preferential partitioning of H₂O into the melt phase over NAMs in (residual) eclogite^{1,2,4}. As these hydrous melts react with enclosing peridotite, H₂O will be in part be incorporated into NAMs in the newly formed orthopyroxene-rich RZLs. However, an orthopyroxene-rich RZL containing 75% orthopyroxene, 20% garnet, and

5% clinopyroxene may only be able to accommodate ~0.03 wt.% of H₂O (i.e. assuming that orthopyroxene, garnet and clinopyroxene take up maxima of ~320 ppm H₂O, ~200 ppm H₂O and ~980 ppm H₂O respectively under water-saturated conditions^{4,24}), which means the excess H₂O may be present as vapour and thus lower the solidi of the orthopyroxene-rich RZL and adjacent mantle lithologies. New melting may then initiate in the orthopyroxene-rich RZL, or olivine orthopyroxenite/websterite formed at the contact between peridotite and orthopyroxene-rich RZL¹³, or continue in the residual eclogite, i.e. depending on the solidus of the particular mantle lithology + H₂O and T_p (Fig. 3).

Reaction and freezing processes may initially only occur at the eclogite-peridotite boundary for incipient Res2 melts at 5 GPa as segregation of melt from the eclogitic residue will be limited by poor interconnection of melt networks/pools (SFig. 3). In contrast, higher Res2 melt fractions at 3 GPa must separate from residual eclogite, as intergranular melt will be strongly interconnected at ~30% melting (SFig. 3). It is conceivable that an original eclogite body (such as Res2) could become strongly zoned during adiabatic ascent due to reaction at the eclogite/peridotite interface. The core of such bodies may comprise a solid residue of Res2 (RRes2) with a higher solidus temperature than Res2, with or without partial melt. Provided that the solidus temperature of the RZL remains above ambient mantle conditions, it may act as a seal to prevent further melt migration and reaction with the surrounding peridotite. In this case, eclogitic partial melts produced by continued adiabatic upwelling will be retained in the eclogite, so that melt and the solid residue remain in equilibrium. Convective motions are likely to cause stirring, stretching and internal deformation of Res2 zoned fragments. Eclogitic bodies may be deformed into layers, even at the cm-scale¹⁸, increasing the reaction surface, leading to opening of the sealing RZL and RRes2 layers, and release of Res2 melt, which freezes into adjacent peridotite on contacts. Continuous mantle convection and stirring may repeatedly re-open and re-seal such rheologically contrasting, refertilised Res2 melt bodies until all melt available has reacted into peridotite, ensuring a wide distribution of frozen 'crustal components' within the Earth's upper mantle.

Residual bimineralic eclogite formed at 3 GPa may become sufficiently Mg-rich and clinopyroxene low in both Ca-eskolaite and jadeite components (Fig. 2; SFig. 7–9) that solidus temperatures are no longer below those of (refertilized) harzburgite/lherzolite, olivine orthopyroxenite/websterite (i.e. the contact between peridotite and orthopyroxene-rich RZL), or orthopyroxenite (RZL) lithologies (Fig. 3). Subsequent melting will then proceed at the melting minimum of olivine orthopyroxenite/websterite \pm H₂O¹³ (TZ in Fig. 3). Melt compositions produced will be nepheline-normative, lying to the silica-undersaturated side of the thermal divide^{13,19,25} in Fig. 3. Due to the relatively high Na/Ca and low Ca/Al ratios of the refertilising eclogitic (Res2) melts (SFig. 5), these new melts will also be characterised by higher Na/Ca and lower Ca/Al ratios ('crustal signature') relative to melt compositions known from nepheline-normative melts of 'pristine' peridotite^{10,26}.

Subduction and subsequent upwelling of residual eclogite Res2 enclosed in ambient mantle may continuously cause numerous other heterogeneities (Fig. 3). For instance, reduced mantle at high pressures²⁷ may impose its low oxygen fugacity (fO₂) on widely distributed bodies of Res2, assuming that recycled crustal material is predominantly oxidized²⁸. This may cause oxidation of CH₄ present in reduced ambient mantle to form H₂O and C (graphite/diamond), and instigate redox melting (reaction 4 in SNotes)^{26,28,29}.

As the adiabat under consideration lies in the (incipient) melting field of peridotite + H_2O under oxidising conditions (Fig. 1b), the presence of significant H_2O may also lead to (incipient) melting of peridotite³⁰. The resulting melts are olivine-rich basanite to nephelinite, enriched in alkalis, with high Ca/Al and Mg[#] ~ 70–75. Reaction of these melts with eclogite such as Res2, will increase the

modal clinopyroxene content and jadeite components (with possibly slightly higher kosmochlor solid solution) as well as decrease the Caeskolaite component of clinopyroxene with little or no change in Mg[#] or Cr/Al of the eclogite. Clinopyroxene enriched zones will be produced at the peridotite/eclogite interface by such 'back-reaction' varying from possible olivine-bearing garnet pyroxenite with little or no orthopyroxene and Mg[#] ~85–90 (metasomatised peridotite) to garnet-bearing clinopyroxenite with Mg[#] ~75–80 and less orthopyroxene (metasomatised eclogite) than in the analogous reaction of siliceous eclogitic melt into peridotite (Fig. 3).

Under oxidising conditions, the presence of significant H_2O may also lead to melting of peridotite + CO_2 + H_2O at greater mantle depths than Res2 along the given adiabat (Fig. 1b). Resulting melts are carbonatitic to carbonate-bearing silicate melts^{26,29,31,32}. If residual eclogite Res2 is swamped by such melts, it will react to form new clinopyroxene-rich RZL (with increased diopside and Ca-Tschermaks solid solution, and reduced or no Ca-Eskolaite component) in the presence of C (graphite/diamond) and/or carbonate and/ or CO_2 and/or SiO₂ (reactions 3–4; Fig. 3; SFig. 11).

$$2Ca_{0.5}[\blacksquare]_{0.5}AlSi_2O_6 + CaMg(CO_3)_2 =$$

$$CaMgSi_2O_6 + CaAlAlSiO_6 + SiO_2 + 2C + 2O_2$$
(3)

Ca-eskolaite (eclogite)

+ dolomite (carbonatitic to carbonate – bearing silicate melts)

= diopside (RZL) + Ca - tschermaks (RZL)

+SiO₂(melt/quartz/coesite; RZL)

+ graphite/diamond/CO₂ (melt/vapour)(RZL)

$$2Ca_{0.5}[\blacksquare]_{0.5}AlSi_2O_6 + MgCO_3 + =$$

$$Mg_2Si_2O_6 + CaAlAlSiO_6 + SiO_2 + 2C + 2O_2$$
(4)

Ca-eskolaite (eclogite)

+ magnesite (carbonatitic to carbonate – bearing silicate melts)

= enstatite (RZL) + Ca - tschermaks (RZL)

+SiO₂ (melt/quartz/coesite; RZL)

+ graphite/diamond/CO₂ (melt/vapour) (RZL).

Thus, reaction with carbonate-rich melts may also cause melting of Res2, producing (sodic) carbonate-bearing silicate or silicatebearing carbonatitic melts (as observed previously^{33,34}), buffered by these reactions. Subsequent reactions of eclogitic carbonate melts with ambient mantle may in turn leach out Cr and Na from peridotitic garnet, possibly producing Cr- and Na-poor garnets enclosed in some diamonds (reactions 5–6 in SNotes), as described above (reaction 2; and reaction 3 in SNotes).

Redox melting and -freezing may thus lead to the co-existence of carbonate- and/or diamond-bearing peridotite and eclogite depending on the pressure, temperature and fO_2 conditions relative to silica-carbonate-diamond redox equilibrium reactions^{26,29,35–37} (reactions 3–4; reactions 4–8 in SNotes). However, there is currently a lack of experimental calibration of these reactions with respect to parameters of pressure, temperature, fO_2 , and composition.

Dynamic mantle processes ensure continuous changes in redox conditions at the interface between eclogites and peridotite, suggesting continuous formation and destruction of diamond and carbonatite along an adiabat. Subduction and recycling of former crust appears to be one of the main causes for continuous but episodic self-driven and -fluxing (redox-) re-melting and -freezing events. Thus, it is not surprising that (incipient) melting and alkaline magmatism occurs episodically and with increased frequency over the Earth's history after commencement of tectonic regimes involving plate subduction³⁸⁻⁴⁰.

Methods

Res2 starting composition was prepared as glassed synthetic sintered oxide powders using established procedures¹³, and doped with a trace-element blend of 10 elements (La, Ce, Sm, Gd, Yb, Sc, Y, Zr, Sr, Cu) at levels of 400 ppm. We used established experimental and analytical procedures as outlined by Spandler et al.¹². The accuracy of the EDS/SEM methods employed has been demonstrated by Spandler et al.⁴¹. Sample preparation, FTIR spectroscopy, and quantification of H₂O and CO₂ in Res2_19 melts were similar to Liu et al.⁴² and Mandeville et al.⁴³. Sample thickness was on average 200 µm. FTIR data, absorbance values, and typical FTIR spectra are given in SFig. 2 and STable 2. Absorbance peaks at ~3550 cm⁻¹ is attributed to O–H stretching modes of H₂O and OH. To determine the H₂O content of the sample, the backgrounds of the corresponding bands were estimated and subtracted from the absorbance by interpolation of a straight line through the minima at the high and low wave-number ends of the absorption peaks. The minute absorbance peak at ~2950 cm⁻¹ represents tiny amounts of organic contamination, and hence, is not further considered.

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Author contributions

The manuscript and the ideas therein were mainly developed by A.R., G.M.Y. and D.H.G., with contributions from all other co-authors. G.M.Y. and D.H.G. initiated and planned the project, which was carried out by A.R. as part of her PhD-project. G.M.Y. and D.H.G. acted as supervisors, and J.H. and C.S. as advisors, and supported A.R. comprehensively during her PhD. A.R. carried out the high pressure experimental work, prepared the experimental charges for SEM analysis, made SEM micro-analyses, compiled all the analyses, formulated results and conclusions as a PhD-thesis, as well as subsequent conclusions such as the new silica-carbonate-diamond redox equilibrium reactions. I.K. supported A.R. substantially in preparing the thin section for FTIR analyses, in performing the FTIR Spectroscopy and in processing of the FTIR data.

Additional information

Supplementary information accompanies this paper at http://www.nature.com/ scientificreports

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