A trace element perspective on the source of ocean island basalts (OIB) and fate of subducted ocean crust (SOC) and mantle lithosphere (SML)

We analyze the first-order observations, basic concepts and explicit/implicit assumptions built into the three major hypotheses for the enriched component(s) in the source of ocean island basalts (OIB) in terms of incompatible trace elements: (1) subducted ocean crust (SOC), (2) subducted continental sediments, and (3) mantle metasomatism. SOC is compositionally too depleted (i.e., [La/Sm]N < 1) to be the major source material for OIB that are highly incompatible element enriched (e.g., [La/Sm]N >> 1). We cannot rule out the contribution of continental sediments as an enriched OIB source component; however, except for two known cases that are yet to be further investigated, there is no convincing evidence for any significant sediment contribution to the petrogenesis of global OIB. Continental materials through subduction erosion can certainly contribute to mantle compositional heterogeneity and may contribute towards some enriched component in OIB source regions. Overall, OIB are not only enriched in incompatible elements, but also enriched in the progressively more incompatible elements, with their inferred source material being variably more enriched than the primitive mantle. These observations require that the OIB sources are pre-enriched by low-degree (Low-F) melt metasomatism. The interface between the growing oceanic lithosphere and the top of the seismic low velocity zone (LVZ) represents a natural peridotite solidus and is the ideal site for major low-F melt induced metasomatism. The ~ 70 Myr history of oceanic lithosphere growth to its full thickness of ~ 90 km records the history of mantle metasomatism, resulting in the deep portion of the oceanic lithosphere being an important enriched geochemical reservoir. We argue that ancient subducted metasomatized mantle lithosphere (SML) provides the major source component for OIB. The metasomatic agent is an H2O-CO2-rich silicate melt derived from within the LVZ. Upward migration and concentration of the melt at the lithosphere-LVZ interface (i.e., the lithosphere-asthenosphere boundary or LAB) results in chemical stratification in the LVZ with the deeper portion being more depleted (i.e., DMM), providing the source for MORB. The widespread metasomatized peridotites, pyroxenites and hornblendites from xenolith suites exhumed from the deep lithosphere (both oceanic and continental) and in orogenic peridotite massifs confirm the role of a low-F silicate melt phase as the metasomatic agent. The SOC, if subducted into the lower mantle, will be too dense to return in bulk to the upper mantle source regions of oceanic basalts, and may have contributed to the two large low shear wave velocity provinces (LLSVPs) at the base of the mantle beneath the Pacific and Africa over Earth’s history.

Introduction

Differences in the geochemical characteristics of mid-ocean ridge basalts (MORB) and intra-plate ocean island basalts (OIB) were already obvious over 40 years ago (Gast, 1968). These differences were instrumental in formulating the mantle plume hypothesis (Morgan, 1971, 1972). OIB are rich in potassium and light rare earth elements, which is consistent with a mantle plume origin from a primordial mantle source deep in the lower mantle, whereas the ocean ridge crests tap only the passively rising shallow asthenosphere that has low potassium and light rare earth element contents because of previous seafloor spreading-related episodes of reworking (Morgan, 1972). This first-order perception remains largely unchanged although our understanding of the mantle geochemistry is much better today.
thanks to the efforts of intensive, world-wide sampling and detailed trace element and isotope studies using analytical methods of varying sophistication over the past forty years.

One of the major advances in this regard is the recognition of mantle compositional heterogeneities through studies of oceanic basalts. OIB are particularly variable in composition such that several isotopically distinct mantle source end-members (e.g., “DMM”, “EM1”, “EM2”, “HIMU”, “FOZO”, “C”) have been proposed to explain the variability (e.g., White, 1985; Zindler and Hart, 1986; Hart et al., 1992; Hanan and Graham, 1996; Hofmann, 1997). The isotopic ratio differences among these end-members reflect the differences of the radioactive parent/radiogenic daughter (P/D) ratios (e.g., Rb/Sr, Sm/Nd, Lu/Hf, U/Pb and Th/Pb) in their ultimate mantle sources which, with time, evolve to distinctive fields in isotope ratio spaces. Significant fractionation of parent/daughter (P/D) ratios in the solid state is considered unlikely in the deep mantle due to extremely slow diffusion rates (Hofmann and Hart, 1978); hence it is logical to suspect that processes known to occur in the upper mantle and crust (e.g., magmatism, metamorphism, weathering and sedimentary processes) are the likely causes of any P/D fractionation. These shallow or near-surface P/D fractionated materials are then introduced into the mantle sources of oceanic basalts through subduction zones. Mantle compositional heterogeneity is thus assumed to be a general consequence of plate tectonics. Despite this apparent conceptual clarity, our understanding of the origin of mantle compositional heterogeneity, in particular the origin of the enriched geochemical signatures of OIB, remains somewhat incomplete.

Here, we do not attempt to solve the OIB source problem, nor to conduct a comprehensive review of the scientific literature, but to analyze the first-order observations, basic concepts and explicit/implicit assumptions built into the major hypotheses for the enriched component(s) in OIB sources in terms of incompatible trace elements. While radiogenic isotopes are important in helping revealing OIB source histories as dealt with by many, we take an incompatible element approach in illustrating the concepts because it is, relatively-speaking, simple, explicit and straightforward relative to that based on radiogenic isotopes; the latter can be quite complex, and their interpretations ultimately rely on an understanding of trace elements, especially the P/D ratios, as discussed above. Willbold and Stracke (2006) have demonstrated that the EM1, EMII and HIMU OIB end-members, defined on the basis of their St-Nd-Pb isotope characteristics, cannot be uniquely distinguished in terms of incompatible trace elements. Because of the wide-spread association of “OIB” with “mantle plumes”, we hope the conclusions reached in this study may contribute to the great mantle plume debate (e.g., Armstrong, 1968; Gast, 1968, Hofmann, 1988), we could be satisfied with the OIB source being less depleted than the MORB source. However, the OIB source is actually more enriched in incompatible elements than the PM (Sun and McDonough, 1989; Niu and O’Hara, 2003, 2009; Prytulak and Elliott, 2007; Humphreys and Niu, 2009). It also varies significantly in its inferred abundances and ratios of incompatible elements, as well as radiogenic isotopes, from one island to another and from one group of islands to another group.

By accepting the notion that mantle compositional heterogeneity results from plate tectonics, it is logical to search for the incompatible element enriched OIB component in this context. Hofmann and White (1982) were the first advocates for “Recycled Oceanic Crust” (ROC) as the enriched source material for OIB. They proposed that oceanic crust is returned to the mantle during subduction. Eventually, it becomes unstable (at the core-mantle boundary; see Christensen and Hofmann, 1994) as a consequence of internal heating, and the resulting diapirs become the source of ocean island basalts (OIB) and other hot-spot volcanism. While some details are considered conjectural, the principal idea of the ROC model has been widely accepted as being correct (see White, 2010).

Some authors have suggested that subducted terrigenous sediments (upper continental crust (CC) derived material) may be responsible for enriched signatures of OIB in terms of both incompatible trace elements and isotopes (e.g., Chauvel et al., 1992; White and Duncan, 1996; Hofmann, 1997; Jackson et al., 2007). Others have emphasized the importance of mantle metasomatism (Sun and McDonough, 1989; McKenzie and O’Nions, 1995; Halliday et al., 1995; Niu et al., 1996, 1999, 2002, 2011; Niu and O’Hara, 2003; Donnelly et al., 2004; Workman et al., 2004; Pilet et al., 2005, 2008, 2011; Niu, 2008, 2009; Humphreys and Niu, 2009).

Subducted ocean crust (SOC) is not a major source for OIB

Niu and O’Hara (2003) showed that OIB mantle source regions and the basic underpinning concepts.

Subducted ocean crust (SOC) is not a major source for OIB

Niu and O’Hara (2003) showed in terms of straightforward petrology, trace element/isotope geochemistry and mineral physics that ancient subducted ocean crust (SOC) cannot be the major source material for OIB. Among many other reasons, SOC, if subducted to the lower mantle, will not return in bulk to the upper mantle, but will sink irreversibly to the base of the mantle. This is because ocean crust of MORB composition is denser than the ambient mantle at all depths except the depth interval between 660 and 720 km (Ono et al., 2001; Hirose et al., 2005; see discussion by Niu and O’Hara, 2003), and it is not straightforward how such dense material can rise in bulk to OIB source regions in the upper mantle. We do not, however, rule out the possible presence of fragments of SOC entrained in some form of deep mantle upwelling. Consequently, it is not surprising if some OIB do indeed contain traces of the geochemical fingerprint of SOC. Note that we need to make a conceptual distinction here between the terms ROC and SOC: ROC is a description, but ROC is an interpretation that SOC is necessarily returned to OIB source regions.
SOC is too depleted to be the source material for OIB

If SOC cannot return in bulk to the mantle source regions for OIB, the concept of "recycled oceanic crust" (ROC) as OIB source may not be practical. We can readily demonstrate that the "ROC" interpretation for OIB has difficulties. Figure 1 shows this analysis schematically in terms of rare earth element (REE) patterns. Obviously, melting of incompatible element depleted SOC cannot produce incompatible-element highly enriched OIB (process 4 in Fig 1), without even mentioning the petrologic unlikeness of partially melting basaltic rocks (SOC) with MgO < 13 wt% alone to produce picritic melts with MgO > 15 wt% (e.g., primitive Hawaiian OIB) (see Niu and O'Hara, 2003). This again simply means that the major Hawaiian source rock is peridotitic rather than basaltic/eclogitic. Figure 2 compares two independent estimates of average OIB (Sun and McDonough, 1989; Willbold and Strake, 2006) with average N-type MORB (Sun and McDonough, 1989) and the mean composition of the ocean crust (Niu and O’Hara, 2003). It is clear that the ocean crust (or SOC when injected into the mantle) is too depleted in incompatible elements to be a possible source material for OIB, as quantitatively demonstrated in Figure 3 below. Furthermore, even if the ancient SOC were indeed present in the OIB source regions, it is isotopically far too depleted to explain the enriched isotopic properties of OIB (see Figure 1 of Niu and O’Hara, 2003).

Figure 3 demonstrates the argument quantitatively using island-averaged OIB data (see Table 2 of Humphreys and Niu, 2009 for data source, derived from the global GEOROC database: http://georoc.mpch-mainz.gwdg.de/georoc/). Since TiO₂ (we use Ti₇₂, where 72 refers to the TiO₂ content in OIB corrected for fractionation effects to an Mg# value of 0.72) and P₂O₅ (P₇₂) are reliably determined major and minor incompatible element oxides routinely analyzed along with the rest of the major element oxides, and because the REEs are reliable incompatible and moderately incompatible trace elements (vs. other incompatible trace elements in the database; see Humphreys and Niu, 2009 for data discussion), our analysis here is focused on these elements, i.e., Ti₇₂, P₇₂, [La/Sm]₇₉ and [Sm/Yb]₇₉. The panels a, b and c on the left in Figure 3 plot Ti₇₂, P₇₂ and [La/Sm]₇₉ against [Sm/Yb]₇₉. While the data trends (thick grey linear regression lines with arrows) largely point to a control by increasing pressure (P) and decreasing extent (F) of melting from beneath thin lithosphere to...
Figure 3 Panels a-c on the left are based on the 115 island-averaged data given in Table 2 of Humphreys and Niu (2009). The thick arrowed lines are regression lines and point in the direction of increasing pressure and decreasing extent of melting with increasing lithosphere thickness (Humphreys and Niu, 2009; Niu et al., 2011). Using the primitive mantle composition (PM, Sun and McDonough, 1989) as the source (purple square) we show that its partial melts by varying extents of partial melting from vanishingly small (~ 0.01%) up to 10% under both garnet (red) and spinel (blue) facies conditions cannot explain the formation of the highly enriched OIB melts. The arrows (downwards and leftwards) in panel b, which applies to all other panels, indicating increasing extent of melting form 0.1 to 10%. The incongruent melting relationship \[0.083 \text{ olivine} + 0.810 \text{ cpx} + 0.298 \text{ garnet} = 0.190 \text{ opx} + 1.000 \text{ melt}\] with initial modes of 0.53 olivine, 0.27 cpx, 0.04 garnet and 0.16 opx (Walter, 1998) is used for garnet peridotite facies melting (red squares). The incongruent melting relationship \[0.652 \text{ opx} + 0.466 \text{ cpx} + 0.049 \text{ spinel} = 0.167 \text{ olivine} + 1.000 \text{ Melt}\] with initial modes of 0.513 olivine, 0.341 opx, 0.131 cpx and 0.015 spinel (Niu, 1997) is used for spinel peridotite facies melting (blue triangles). The simple batch melting model \((C_1/C_0 = 1/(D_0 + F(1 – P)))\) is adequate to illustrate the concept. Relevant partition coefficients are all taken from Prytulak and Elliot (2007). The panels d-f on the right are the same but also show the model bulk composition of the ocean crust (bluish grey diamond; Niu and O’Hara, 2003) and its partial melts (as indicated) by varying extents of melting from vanishingly small (~ 0.01%) up to 10% under eclogite facies conditions using modes of 0.6 omphacitic cpx and 0.4 garnet appropriate for MORB protoliths (Song et al., 2006). For conceptual illustration, we did not include rutile and assume modal batch melting with the bulk ocean crust approximates the subducted oceanic crust (SOC) for the elements concerned, these simple calculations allow effective evaluation of the actual roles of SOC in the genesis of OIB melts. Note that inclusion of rutile will lower the TiO\(_2\) in the model melt in the Ti\(_{72}/\text{Sm/Yb}\)\(_{PM}\) (see text for details).
beneath thick lithosphere (see Figure 1 of Niu et al. 2011), the large compositional range and the scatter cannot be produced by melting a uniform mantle source. To illustrate the argument, we use the PM composition (Sun and McDonough, 1989) as the source to calculate melt compositions generated by ~0.01% (vanishingly small) to 10% batch melting (simple and adequate for this purpose) under both spinel and garnet peridotite facies conditions as indicated (see the caption to Figure 3 for further details). As expected, a variety of sources varying from PM-like to variably much more enriched than PM would be required to explain the OIB compositional spectrum, in particularly, the data with Ti$_{pm}$, P$_{pm}$, [La/Sm]$_{pm}$ and [Sm/Yb]$_{pm}$ values greater than can be modelled with the extent of melting vanishingly small (Figure 3).

The panels d, e and f on the right in Figure 3 include the bulk ocean crust composition (Niu and O’Hara, 2003) and its partial melts from ~0.01 to 10% batch melting under eclogite facies conditions (see Figure 3 caption for details). Assuming the bulk ocean crust approximates the SOC composition, these simple calculations suggest that SOC, if present at all in OIB source regions, contributes very little to the enriched OIB compositions. In the Ti$_{pm}$-[Sm/Yb]$_{pm}$ Plot (Figure 3d), it seems possible that SOC melts could represent the enriched end-member for OIB melts. However, such SOC melts cannot contribute in any straightforward way to OIB compositions in the P$_{pm}$-[Sm/Yb]$_{pm}$ (Figure 3e) and [La/Sm]$_{pm}$-[Sm/Yb]$_{pm}$ (Figure 3f) plots. Furthermore, the proportion of SOC is likely volumetrically small and would be preferentially melted out because of its low solidus temperature and heat-suction effect during melting (Hirschmann and Stolper, 1996; Stolper and Asimow, 2007). As a result, the contribution of SOC by total melting to OIB would be the bulk composition of the SOC itself, which lies at the most depletred end of the OIB compositional spectrum. Therefore, SOC contributes, if not actually diluting, very little to the enriched component of OIB.

Revised “ROC” model has more difficulties

In recent years, the ROC model has become more popular because of some refreshed lines of evidence (e.g., Sobolev et al., 2000, 2005, 2007). Sobolev et al. (2000) interpreted the composition of olivine melt inclusions in some Hawaiian lavas as recording the signature of “ghost plagioclase” from ancient recycled gabros (lower ocean crust). This interpretation has gained some acceptance, but as the host olivine of these melt inclusions is a liquidus phase crystallized from cooling and evolving melts in shallow crustal magma chambers, the trapped melt cannot be a primary melt in equilibrium with mantle minerals and evolving melts in shallow crustal magma chambers, the trapped melt cannot be a primary melt in equilibrium with mantle minerals (Niu and O’Hara, 2003). Thus, it is unclear how the composition of the melt inclusions can be used to infer Hawaiian OIB source lithologies. Furthermore, it is intriguing as to why such a source-indicating “ghost plagioclase” signature should be preserved only in these minute melt inclusions, yet not in the host lavas. This conundrum was answered elegantly by Danyushhevsky et al. (2004), who demonstrated that Al-Ca-Sr rich melt inclusions in phenocryst olivines result from a complex grain-scale dissolution-reaction-mixing process, and that dissolution of phenocryst plagioclase can leave such a signature in the melt inclusions and does not in any way demonstrate a role for recycled gabros in the petrogenesis of Hawaiian basalt.

Sobolev et al. (2005) argued that the high Ni content of olivine phenocrysts in some Hawaiian lavas reflects the presence of volumetrically significant “SOC” in the mantle source region. The “SOC”, in the form of eclogite, melts first to produce SiO$_2$-saturated melts which then react with surrounding harzburgite to produce an olivine-free pyroxenite in the Hawaiian magma source region. The absence of olivine in the resultant source region could explain the high Ni content in the parental melts and hence the high Ni contents of the olivine phenocrysts crystallized from Ni-rich basalts in shallow-level magma chambers beneath Hawaii. Such a conceptually (not necessarily geologically and physically) feasible process was invoked to emphasize the requirement of recycled “SOC” in the petrogenesis of Hawaiian OIB in favour of the “ROC” model. However, it has been demonstrated in terms of straightforward phase equilibria that Hawaiian basalts require olivine to be present in the magma source region (Green and Falloon, 2005; Presnall and Gudfinnson, 2009, 2011). Furthermore, a robust Ni-olivine Kd and mass balance analysis (Li et al., 2008, Li and Ripley, 2010) has pointed out that the calculations and interpretations by Sobolev et al. (2005) on Hawaiian lavas and olivines are unsupported. More recent experimental studies on $K_{Olivine/melt}$ as a function of $T$ (Wang and Gaetani, 2008; Matzen et al., 2009) demonstrate that the high Ni olivine in Hawaiian melts can be readily produced without having to invoke an olivine-free pyroxenite source. We consider that while an eclogite-derived melt reaction with harzburgite is a petrologically interesting concept, it may actually not take place at all in practice because of the physical difficulty in transporting volumetrically significant amounts of dense eclogite into the source region of Hawaiian magmatism. Importantly, the invoked eclogite of the “SOC” protolith is highly depleted in incompatible elements (Figs. 1 and 2), as is the harzburgite with which the eclogite melt react (unless metasomatized; see below). Therefore, this revised ROC (R-ROC) model cannot produce OIB that are highly enriched in incompatible elements (Figs. 2 and 3).

Sobolev et al. (2007) have gone a step further to argue that the Ni, Cr, Mn and Ca contents of olivine phenocrysts can be used to estimate the proportions of “SOC” in the mantle source regions of basalts erupted both in the ocean basins and on the continents. They showed that olivine Ni contents are high in basalts erupted on thick (~70 km) lithosphere, low in basalts erupted on thin (<70 km) lithosphere, and lowest in MORB. This, plus the correlated variations of Cr, Mn and Ca in olivine, allowed them to quantify that “SOC” in OIB source regions is necessarily more abundant beneath thick lithosphere than beneath thin lithosphere. They stated that the proportion of “SOC-eclogite” in mantle source regions increases with increasing lithosphere thickness (e.g., 5% beneath ocean ridges, 10% in Iceland mantle, and 20% in Hawaiian mantle). As a result, they proposed that “SOC eclogite induced olivine-free pyroxenite” contributes more to the erupted basalts with increasing lithosphere thickness: ~10-20% for MORB, ~20% for Iceland “OIB”, ~40% for Detroit seamount OIB, ~60% for Hawaiian OIB and ~100% for the Siberian flood basalts.

We have demonstrated that the high quality compositional data provided by Sobolev et al. (2007) for olivines in MORB and basalts erupted both on thin and thick lithosphere in the ocean basins and on the continents are wholly consistent with and readily explained by the variation in the lithospheric thickness or the lid effect (Niu and O’Hara, 2007; Niu et al. 2011) without the need to invoke varying abundances of SOC in the mantle source regions, which has more difficulties. It is intuitively unreasonable that SOC should be more abundant beneath old thick lithosphere than beneath young thin lithosphere. If SOC were passively embedded in a peridotite matrix in the upper mantle, there would be no straightforward physical mechanism that selectively drives the SOC material from beneath
thin lithosphere to beneath thick lithosphere to be sampled by OIB volcanism. If SOC was incorporated into hot mantle plumes, it is fortuitous that deep-rooted plumes would know the “required” amount of SOC as a function of lithosphere thickness near the Earth’s surface. As SOC has a lower melting temperature than the ambient peridotite mantle, it is possible that SOC might contribute more to OIB melts because of the restricted extent of melting beneath thick lithosphere, but SOC is compositionally too depleted to yield highly enriched OIB (Figures 1-3).

Subducted continental sediment is not a major source for OIB

Many authors have suggested that subducted terrigenous sediments could be responsible for the incompatible element enriched signature of some OIB (see discussion above). In the context of plate tectonics with mantle circulation, it is possible and also likely that terrigenous sediments can be transported into the source regions of oceanic basalts. The question remains as to whether these sediments are indeed the enriched component seen in OIB. Looking at the REE patterns (Figure 4a), the similarity of average OIB (Sun and McDonough, 1989; Wilbold and Strake, 2006) to bulk continental crust (CC; Rudnick and Gao, 2003) and global subducting sediments (GLOSS; Plank and Langmuir et al., 1998) is encouraging, but we can readily see the difficulties when other incompatible trace elements are considered (Figure 4b). The CC and GLOSS have characteristic depletions in Nb, Ta, P and Ti and enrichment in Pb - such a “CC-like signature” is rather strong and should be evident in the incompatible element characteristics of OIB if terrigenous sediments were indeed the enriched component. In fact, if anything, OIB are relatively enriched, not depleted, in Nb (vs. Th) and Ta (vs. U) (Figure 4b; also see Niu and Batiza, 1997).

Thus far, the best evidence for OIB suites with an apparent “CC-like signature” is from the Naturaliste oceanic plateau in the Indian Ocean (Mahoney et al., 1995) and some lava flows from Samoa (Savai) in the Southwest Pacific (Jackson et al., 2007). As the former is geologically close to a passive continental margin with Archean crust whereas the latter is close to the active Tonga subduction zone with a forearc possibly associated with ancient continental lithosphere (Niu et al., 2003), it is unclear if the CC-like signatures in these two cases are indeed produced by recycled terrigenous sediments or continental lithosphere material made available at shallow levels. Further studies of these two cases, based on a combined approach involving petrology, geochemistry, geology and geophysics should prove useful.

In this context, it is necessary to briefly discuss the recent work by Rapp et al. (2008), who conducted ultrahigh pressure (16 to 23 GPa) experiments on terrigenous sediments and advocated the significance of their findings in explaining some mantle compositional heterogeneities. They discovered that the stable high-P assemblage contains 15 to 30% K-hollandite, which controls much of the whole-rock budget of large-ion lithophile elements (LILE, e.g., Rb, Ba, Sr, K, Pb, La, Ce and Th). They concluded that the incompatible trace-element signature of EMI OIB sources can be attributed to recycling of K-hollandite-bearing continental sediments to Transition Zone depths. We note, however, that the geochemical characteristics of the sediments remain unchanged (Figure 5), and differ distinctly from OIB compositions.

In the context of plate tectonics with sediment subduction (e.g.,

Continental mantle lithosphere may be a potential enriched source for OIB

McKenzie and O’Nions (1995) suggested that subcontinental lithospheric mantle (SCLM) could be an important source component for OIB. Because the SCLM is compositionally depleted (in terms of a basaltic component) and therefore physically buoyant, the
Mantle metasomatism of oceanic lithosphere can explain OIB source enrichment

Why mantle metasomatism?

It has been recognized that OIB source materials are enriched in incompatible elements relative to the PM, e.g., \([\text{La}/\text{Sm}]_{\text{OIB Source}} > [\text{La}/\text{Sm}]_{\text{PM}}\) (e.g., Sun and McDonough, 1989; McKenzie and O’Nions, 1995; Niu et al., 2002; Niu and O’Hara, 2003; Prytulak and Elliott, 2007; Humphreys and Niu, 2009; Figure 3), and that OIB are more enriched in the progressively more incompatible elements (Figure 6). These observations require that OIB sources have undergone prior enrichments through a process of low-degree (low-\(F\)) melt infiltration, which is termed mantle metasomatism (e.g., Sun and Hanson, 1975; Lloyd and Bailey, 1975; O’Reilly and Griffin, 1988; Pilet et al., 2005). Wyllie and his collaborators made pioneering contributions to our understanding of mantle metasomatism through experimental petrology and its application to global tectonics and the Earth’s chemical differentiation (e.g., Wyllie, 1984, 1985, 1987a,b,c, 1988a,b; Wyllie and Huang, 1975,1976; Wyllie and Sekine, 1982; Sekine and Wyllie, 1982a, b; Wyllie et al., 1983; Gaspar and Wyllie, 1987; White and Wyllie, 1992; Baker and Wyllie, 1992; Wyllie and Ryabchikov, 2000).

The pre-metasomatic sources may be primitive mantle or previously depleted melting residues, and their metasomatism has been ascribed to infiltration by a low-\(F\) melts that are enriched in volatiles (e.g., \(H_2O\), \(CO_2\)) and incompatible elements. The evidence for metasomatism has come from mantle xenoliths in oceanic and continental alkali basalts and kimberlites (e.g., Frey and Green, 1974; Frey et al., 1978; Menzies and Murphy, 1980; Menzies, 1983; Menzies and Hawkesworth, 1987; O’Reilly and Griffin, 1988; Coltorti et al., 1999; Gregoire et al., 2000; Sen et al., 2005; Coltorti and Gregoire, 2008).
and tectonically exhumed massif peridotites (e.g., Frey et al., 1985; Takazawa et al., 2000) as well as inferences from highly enriched mantle derived melts (e.g., Sun and Hanson, 1975; Lloyd and Bailey, 1975).

**Locations of mantle metasomatism**

Wyllie (1980, 1987a, 1988a,b) presented elegant metasomatic models for the petrogenesis of highly enriched magma types such as kimberlite and carbonatite in continental settings and basanite and nepheline in the ocean basins. He stressed the importance of incipient partial melting in the presence of both H$_2$O and CO$_2$ in generating the metasomatic agents which can modify both oceanic and continental lithosphere (e.g. White and Wyllie, 1992; Baker and Wyllie, 1992).

Compared to MORB, OIB samples from all the intra-oceanic islands so far studied are enriched in incompatible elements with [La/Sm]$_{PM}$ $\geq$ 1. Furthermore, many intra-oceanic seamounts on old ocean crust apparently unrelated to "plumes" or "hotspots" (e.g., Batiza, 1982; Batiza and Vanko, 1984; Castillo et al., 2010), including the very young (~ 6 Ma) "Petit Spots" alkali basalts erupted on the 135 Ma old Pacific plate (Hirano et al., 2006), are also highly enriched in the incompatible elements.

All these suggest that mantle metasomatism is widespread in the oceanic lithosphere, and is likely also true in the mantle lithosphere beneath continents. In principle, mantle metasomatism cannot take place in regions of major melting such as beneath ocean ridges and in mantle wedges above subduction zones because of high-F (vs. low-F) melting where less enriched or diluted melts prevail. However, because melting regions are finite, low-F melt may exist and metasomatism may thus take place in the peripheral areas of these major melting regions (Niu et al., 1996; Pilet et al., 2011). Mantle wedge overliving subduction zones is surely metasomatized (see Wyllie and Sekine, 1982; Donnelly et al., 2004), but the connection to OIB petrogenesis is obscured because the metasomatic agent there may have an arc-melt signature (i.e., [Nb/Th]$_{PM}$ (IAB) $\leq$ 1 and [Ta/U]$_{PM}$ (IAB) $\leq$ 1), whereas both MORB and OIB all have [Nb/Th]$_{PM}$ (MORB, OIB) $\geq$ 1 and [Ta/U]$_{PM}$ (MORB, OIB) $\geq$ 1 (Niu and Batiza, 1997; Niu et al., 1999; Niu and O’Hara, 2009; Sun and McDonough, 1989; Halliday et al., 1995; Hofmann, 1997).

The interface between the base of the growing oceanic lithosphere and the seismic low velocity zone (LVZ), which is also termed lithosphere-asthenosphere boundary (LAB), atop the asthenosphere as the ideal site for mantle metasomatism is relevant to OIB petrogenesis (Halliday et al., 1995; Niu et al., 1996, 2002, 2011; Niu and O’Hara, 2003, 2009; Niu, 2008).

**Mechanism of mantle metasomatism at the growing oceanic lithosphere-LVZ interface (LAB)**

Figure 7a shows that oceanic lithosphere grows with time through basalt accretion of LVZ material (red arrows) before reaching its full thickness after ~70 million years. The presence of an incipient melt due to the effect of volatiles (H$_2$O+CO$_2$) in lowering the peridotite solidus is required to explain the characteristics of the LVZ (Lambert and Wyllie, 1968, 1970; Anderson and Spetzler, 1970; Green, 1971, 1991; Wyllie, 1971; Wyllie and Huang, 1975,1976; Green and Lieberman, 1976; Wyllie, 1977; Niu and O’Hara, 2003; Green and Falloon, 2005; Mierdel et al., 2007; Niu, 2008; Green et al., 2010). This incipient melt is enriched in volatiles (e.g., H$_2$O, CO$_2$) and incompatible elements (Niu et al., 2002; Niu and O’Hara, 2003). As the melt is buoyant, it tends to concentrate into a “melt-rich layer” (in green) atop the LVZ, leading to compositional stratification within the LVZ. The deep portion of the LVZ (in yellow) is thus depleted (see below). During the process of lithosphere growth, spinel-garnet lherzolite in the uppermost LVZ is thermally captured as newly accreted lithosphere. The low-F melts (from the melt-rich layer) collect and ascend, crystallizing hydrous minerals within the ambient peridotite (modal metasomatism; O’Reilly and Griffin, 1988; Wilshire, 1987; Nielson and Noller, 1987; Harte et al., 1993; Nielson et al., 1993; Nielson and Wilshire, 1993; Pilet et al., 2008, 2011), forming veins of garnet pyroxenite, hornblende-pyroxenite and hornblendeidite (yellow veins) before being completely absorbed into the ambient peridotite (cryptic metasomatism; O’Reilly and Griffin, 1988; Pilet et al., 2011).

Figure 7b presents a parcel of mantle (diapir or “plume”) that ascends and partially melts by decompression when it intersects the solidus. As these “plume” melts approach the base of the lithosphere, they may gain additional incompatible element enrichments from the melt-rich layer (green). “Plume” melts ascending further through the lithosphere can assimilate earlier-formed metasomatic veins, leading to further enrichment of the OIB melts that are ultimately erupted (Pilet et al., 2008, 2011; Niu, 2008). The erupted OIB melts may thus have three components: (I) fertile mantle source materials from greater depths (plumes?), (II) the LVZ melt layer (green), and (III) components assimilated from the earlier-formed metasomatic vein lithologies in the oceanic lithosphere. Note also that for the petrogenesis of OIB the fertile materials from depth (i.e., component “I”) must be volumetrically dominant, compositionally peridotitic, and may contain recycled, previously (ancient) metasomatized, deep portions of oceanic lithosphere that are already enriched in the volatiles and incompatible elements (Niu and O’Hara, 2003). Also, component “I” is required for OIB petrogenesis to explain the basaltic-picritic composition and the first-order coupling between incompatible elements and radiogenic isotopes in many OIB suites (see Niu and O’Hara, 2003; Niu et al., 2011). For example, the highly scattered variation of $^{87}$Sr/$^{86}$Sr, $^{143}$Nd/$^{144}$Nd, $^{206}$Pb/$^{204}$Pb and $^{176}$Hf/$^{177}$Hf in average OIB as a function lithosphere thickness (Niu et al., 2011) demonstrates OIB compositional heterogeneity, but also shows weak, yet expected trends reflecting coupling of the isotopic compositions with radioactive parent/radiogenic daughter ratios (P/D; e.g., Rb/Sr, Sm/Nd, U/Pb, Th/Pb and Lu/Hf). Such coupling requires a time interval between the formation of the metasomatized lithosphere and the volcanism in excess of 1.0 Gyrs, which may represent the minimum period from the time of the subduction of ancient oceanic lithosphere into the deep mantle to the return of these materials to the OIB source regions (Niu and O’Hara, 2003). Components “II” and “III” (Figure 7b) are not required for contemporaneous OIB volcanism, but their presence and involvement, if any, can contribute to the petrogenesis of extremely enriched magma compositions such as alkali basalt, basanite and nepheline on many ocean islands (e.g., Pilet et al., 2008; Niu, 2008) and some intraplate seamounts (e.g., Batiza and Vanko, 1984). If these components were involved, then incompatible elements and radiogenic isotopes would be decoupled because of significant parent-daughter element fractionation by the recent metasomatism (Figure 7), that is too young/recent to produce significant radiogenic isotope ingrowths. The association of incompatible element enrichment with a depleted radiogenic isotopic
lithosphere metasomatism is a widespread phenomenon, and the deep portion of the oceanic lithosphere is therefore an important enriched geochemical reservoir (Niu and O’Hara, 2003).

The nature of the metasomatic agents

The observation that metasomatic vein lithologies (e.g., pyroxenite, hornblende, other amphibole- and phlogopite-bearing assemblages, as well as grain-boundary enrichment) are common in deep portions of both oceanic and continental lithosphere strongly suggest that the metasomatic agents originate from the seismic LVZ. The observation that OIB are more enriched in the progressively more incompatible elements with respect to major mantle minerals (Figure 6) requires that the metasomatic agents are very low-$F_{H_2O}$ and $CO_2$-rich silicate melts. In recent years, carbonatite melt has been a popular choice of metasomatic agent. Wyllie was among the very first to recognize that the presence of $CO_2$ in the LVZ can induce melting more effectively than $H_2O$ although he considered that both $CO_2$ and $H_2O$ must be at work in the LVZ (Wyllie and Huang, 1975, 1976; Wyllie, 1977; Wyllie and Lee, 1998; Wyllie and Ryabchikov, 2000). Several investigators have proposed recently that the LVZ may be largely or perhaps entirely caused by $CO_2$-bearing/generated incipient melts (e.g., Yaxley et al., 1998; Presnall and Gudfinnsson, 2005, 2008; Dasgupta et al., 2007). These arguments are based on the recognition that the solubility of $H_2O$ in mantle minerals is high enough to absorb any free $H_2O$ (e.g., Hirh and Kohlstedt, 1995; Karato and Jung, 1998), yet the solubility of $CO_2$ in upper mantle minerals is “diminishingly low” (Keppler et al., 2003). It follows that the free volatile components available in the upper mantle must be $CO_2$ or dominated by $CO_2$ in non-supra-subduction zone environments. This naturally leads to the conclusion that $CO_2$-rich melt or carbonatite melt may in fact be the metasomatic agent in the mantle. This argument is further strengthened by the fact that carbonate melts are much more mobile and can readily metasomatize the lithosphere on large scales because of their low viscosity and low solidus temperature.

There is indeed much observational evidence from studies of mantle xenoliths for the importance of carbonatite metasomatism (e.g., Ionov, 1998; Schleicher et al., 1998; Harmer and Gittins, 1998; Coltorti et al., 1999; Gregoire et al., 2000; Newmann et al., 2004). Nevertheless, we consider that $H_2O-CO_2$-rich silicate melts are far more important as metasomatic agents than carbonatite melt alone.

**Figure 7** Oceanic lithosphere thickens with time by basal accretion of LVZ material (red arrows) for ~70 million years before reaching its full thickness of ~90 km in terms of the plate model. The thick purple curve is the present-day interface (LAB) between the growing lithosphere and the LVZ, which is conceptualized as a natural solidus marking the petrologic transition from solid amphibole/phlogopite-bearing peridotite (lithosphere) to peridotite containing a small melt fraction (LVZ). The thin white dashed curves indicate where this interface was in the past, illustrating the continuing lithosphere growth with time as the plate ages. An incipient melt may form at depth (as deep as ~180 km?) in the LVZ caused by the presence of H-C-O volatiles. The incipient melt is thus enriched in these volatiles (e.g., $H_2O$, $CO_2$) and incompatible elements. As the melt is buoyant, it tends to concentrate towards the top of the LVZ as indicated by the green “melt layer” (the thickness is exaggerated for illustration), leading to compositional stratification within the LVZ. The dashed-blue lines with arrows point in the direction of asthenospheric flow because of ridge suction (Niu & Hekinian, 2004). As a result, MORB samples the more depleted deep portions of the LVZ whereas the enriched “melt layer” atop the LVZ continues to contribute to lithosphere metasomatism before it reaches its full thickness. In the process of the lithosphere thickening, the melt from the “melt layer” will collect and ascend, crystallizing hydrous minerals within the ambient peridotite (i.e., modal metasomatism), forming veins of pyroxenite and hornblende (yellow veins) before being finally/completely absorbed in the ambient peridotite (i.e., cryptic metasomatism). Because it takes ~70 million years for the lithosphere to reach its full thickness, metasomatism continues for this length of time. [b] A parcel of mantle material (perhaps a “plume”?) ascends and partially melts by decompression. These “plume” melts (I) may gain additional enrichments from the “melt layer” (II). Continued ascent of the “plume” melts through the lithosphere can cause assimilation of earlier-formed metasomatic veins (III), leading to further enrichment of the ultimately erupted OIB melts. Note that while the “asthenosphere” is often considered synonymous with the LVZ, we here consider the LVZ as the upper portion of the asthenosphere because the latter is present throughout the upper mantle, yet the LVZ is seismically detected mostly beneath ocean basins only. Modified from Niu and O’Hara (2003, 2009) and Humphreys and Niu (2009).
for several reasons. (1) Most metasomatic lithologies so far documented from the continental and oceanic lithosphere are dominated by pyroxenite and hornblende veins (plus garnet, phlogopite, rutile, zircon, titanite), with or without minor carbonate. (2) OIB are enriched in the progressively more incompatible elements, which is consistent with the metasomatic agent being a low-$F_{\text{H}_2\text{O}}$-$\text{CO}_2$-rich silicate melt, rather than a carbonatite melt. Natural carbonatites have trace element systematics that are distinct from those of OIB (Figure 8a). High pressure experimental simulation demonstrates that behaviours of incompatible elements in carbonatite melts differ remarkably from their behaviours in OIB (Figure 8b). Carbonatite melts of deep transition-zone origin also show distinct incompatible trace element systematics (Figure 8c).

Therefore, $\text{H}_2\text{O}-\text{CO}_2$-rich silicate melt, not carbonatite melt, is likely the major metasomatic agent for the petrogenesis of OIB even though carbonatite melt may be locally important, particularly in the continental settings.

Some new insights on mantle dynamics

The above discussion of OIB sources offers some new insights into a number of fundamental aspects of mantle geodynamics.

A new view on the origin and nature of the depleted MORB mantle

The current paradigm of solid earth geochemistry is that the upper mantle is depleted in incompatible elements (e.g., Hofmann, 1997), and is therefore termed depleted mantle (DM) or depleted MORB mantle (DMM) (Zindler and Hart, 1986; see Figure 1). The logic behind this is reasonable because MORB are depleted in incompatible elements and the MORB source must therefore be depleted. Because MORB are produced by decompression melting of passive (vs. dynamic) upwelling asthenosphere, the maximum depth of melting must be shallow. Therefore, the DM or DMM reservoir must also be shallow and may occupy the entire upper mantle or upper part of the lower mantle (see Donnelly et al., 2004; Workman and Hart, 2005; Niu and O’Hara, 2009). The requirement for the DMM to produce highly incompatible element enriched, near solidus (low-$F$) melts that metasomatize the base of the overlying oceanic lithosphere may therefore seem contradictory. This apparent contradiction does not negate the MORB source being depleted, but rather suggests that the standard concept for the DM or DMM (see Figure 1) needs some revision as pointed out by Niu and O’Hara (2009). In Figure 7, the blue dashed lines with arrows point in the

patterns do not favour such carbonatite melt as a metasomatic agent for the enriched signature of OIB. Also, the experimental data show Nb and Ta depletion, but no Ti depletion, yet Zr-Hf enrichment. [c] 3 calculated carbonatite melts in equilibrium with majorite and perovskite stable under Transition Zone conditions (Walter et al., 2008). Carbonatites with such trace element systematics may indeed exist in deep parts of the upper mantle or Transition Zone, but if these were metasomatic agents, they would impart such signatures to OIB, but this is again not observed. Therefore, these three cases suggest that even though carbonatitic melts may indeed exist and may be an important metasomatic agent in some mantle environments, they cannot be the major metasomatic agent required for the enriched signature of the global OIB.
direction of the asthenospheric flow required to feed the ocean ridge because of ridge suction (Niu & Hekinian, 2004). As a result, MORB sample the deep portions of the LVZ (in yellow) that are already depleted by the removal of a low-F’ melt. This concept offers an entirely new perspective on the origin and nature of the depleted mantle (Niu and O’Hara, 2009).

In summary, the compositional stratification in the LVZ (Figure 7) is required by the observations: (1) the LVZ is most consistent with the presence of a volatile-rich melt phase; (2) a melt-rich layer atop the LVZ (green) is required by the occurrence of metasomatism at the base of the growing oceanic lithosphere, which is also required by the abrupt seismic velocity drop at the lithosphere-LVZ boundary (or LAB; Kawakatsu et al., 2009; see Niu et al., 2011 for discussion); and (3) ridge-ward asthenosphere flow resulting from ridge-suction (Niu & Hekinian, 2004) to feed MORB requires that the deep portion of the LVZ is compositionally depleted, i.e., the DMM source.

**Fate of subducted oceanic crust**

We have demonstrated above that SOC cannot be the major source for OIB. The question then is what may have happened to SOC in the context of plate tectonics and mantle circulation over Earth’s history. If ocean crust of MORB composition can indeed sink to the lower mantle as illustrated by global seismic tomography studies (e.g., van der Hilst et al., 1997; Grand et al., 1997; Karason and van der Hilst, 2000), subducted ocean crust (SOC) will not return in bulk to the upper mantle source regions of oceanic basalts because it is denser than the ambient mantle at all depths except the depth interval between 660 and 720 km (see Ono et al., 2001; Niu and O’Hara, 2003; Hirose et al., 2005). If this irreversible process is indeed the case, then this has profound implications for the chemical structure of the mantle as discussed by Niu and O’Hara (2003):

“As discussed above ..., oceanic crust subducted into the lower mantle will not return in bulk to the upper mantle because of the negative buoyancy in both solid and liquid states. Transfer of basaltic crust to the lower mantle would be an irreversible process. This supports the argument for a hidden component deep in the lower mantle that has not been sampled by known volcanism ..., and would also lead to chemical stratification of the mantle with the mean composition of the lower mantle becoming progressively enriched in residual ocean crust lithologies (i.e., compositionally lower in Ca/Al, and higher in Fe/Mg, Si/Mg, Al, and water insoluble incompatible elements such as Ti, Nb, Ta, Zr, and Hf). If subduction of oceanic crusts into the lower mantle has continued for some time, then a large compositional contrast in terms of these elements must exist between the upper and lower mantle.”

We can conduct a simple exercise following this argument. By assuming that (1) plate tectonics began at ~ 4.0 Ga (unknown; see Stern, 2007), (2) the oldest ocean crust may not be older than 200 Myrs before subduction as is the case at present, (3) oceanic crust has always occupied 65% of the Earth’s surface area (vs. 35% area for the continental crust), (4) the average thickness of the ocean crust is about 6 km with a mean density of 3.0 g cm⁻³ (vs. 3.3 g cm⁻³ for the mantle), then the amount of SOC since 4.0 Ga would be ~ 3.0 wt% of the entire mantle. That is, there is 4 Gys/200 Myrs = 20 times the present-day ocean crustal mass stored in the deep mantle. This calculation, based on reasonable and explicit assumptions, is clearly only illustrative. We note that there have been several alternative estimates involving more complex assumptions that give variably greater values. For example, Helffrich & Wood (2001) estimated that ~ 5 wt% SOC is randomly distributed throughout the mantle.

**Is there any evidence for the SOC accumulated in the deep lower mantle?**

We suggest that the two large low shear velocity provinces (LLSVPs) in the lowermost mantle beneath the Pacific and Africa may represent the SOC accumulated over Earth’s history. These two large LLSVPs have been recognized for some time (e.g., Dziewonski, 1984; Grand et al., 1997; Su and Dziewonski, 1997; Ritsema et al., 1999; Kellogg et al., 1999; Mégnin and Romanowicz, 2000). They have been interpreted to be excessively hot mantle domains representing the locations of “superplumes” (e.g., Romanowicz and Gung, 2002; Ni et al., 2002), and to be responsible for the geographically associated surface geoid highs or “superswells” (McNutt, 1988) and probably also mantle isotopic anomalies revealed from volcanism on these highs and their peripheral regions (e.g., Hart, 1984; Castillo, 1988). Recent studies suggest that the LLSVPs have sharp boundaries with, and higher density (~ 2-5%) than, the ambient mantle, indicating that they are chemically distinct from the surrounding mantle (Ni et al., 2002; Becker and Boschi, 2002; Ni and Helfringer, 2003; Wang and Wen, 2004; Toh et al., 2005; Ford et al., 2006; Garnero et al., 2007). Their origin is not well understood and their geodynamic effects cannot yet be properly modelled (e.g., McNamara and Zhong, 2004; Lassak et al., 2010). They could be residual Fe-rich material from core formation or subducted ocean crust (e.g., Garnero et al., 2007; Hirose and Lay, 2008).

Our estimate of ~ 3 wt% SOC in the entire mantle that at present has accumulated over the course of geological time is similar to the estimated mass of the LLSVPs, ~ 2 wt% of the mantle, by Burke et al. (2008). Given the likely uncertainties in these estimates we do not wish to overstate the significance of the calculations, but they do offer an independent line of reasoning in support of the concept that mantle plumes do not originate from recycled oceanic crust and that subducted ocean crust is too dense to return in bulk to the upper mantle source regions of oceanic basalts (Niu and O’Hara, 2003).

**The “buoyant” superswells overlie the dense LLSVPs – a contradiction or natural consequence?**

The geoid highs, superswells, and reduced seismic velocity in the lower mantle are all consistent with mantle “superplumes” initiated close to the CMB or lower mantle beneath the Pacific and Africa. However, because the LLSVPs at the base of the mantle beneath these two regions are not buoyant, but rather dense (~ 2-5% denser than the ambient mantle), it is difficult to see how they could be the source of the “superplumes”. This apparent contradiction points to a serious physical problem – how could this relationship be possible in terms of straightforward physics? Models that apply an imposed plate history can result in focusing of the subducted materials into piles at the core-mantle boundary beneath Africa and the Pacific, corresponding to the two LLSVPs (e.g., Garnero et al., 2007). These models can explain the location of the two LLSVPs, but still cannot explain the observation that the “massive upwellings” manifested by the superswells are underlain by regions of dense materials of huge
negative buoyancy. The “superplume” models suggest that the negative buoyancy of the dense materials can be overcome by thermal buoyancy (cf. Garnero et al., 2007), thus leading to the upwelling of the “superplumes”. Yet, it seems unlikely that thermal buoyancy can be sufficient to overcome the > 2% density contrast (Niu and Batiza, 1991b,c), especially because of the very low thermal expansion coefficients at the deep lower mantle conditions (Anderson, 2007).

Here we suggest that the existence of buoyant superswells overlaying the dense LLSVPs is not a contradiction, but a natural consequence in terms of plate tectonics, petrology and mineral physics. Transformation of the basaltic ocean crust (MORB + gabbroic cumulates) into eclogite during subduction and further high pressure phase transformation at the 410 km seismic discontinuity facilitate subduction. Subducting oceanic lithosphere (or slab) can penetrate the 660 km discontinuity (with some difficulty; see van der Hilst et al., 1997) and enter the lower mantle, sinking eventually to the core-mantle boundary region.

Figure 9 shows that under lower mantle conditions, ocean crust with MORB composition is significantly denser (~ 3.6%) than the ambient mantle (PREM), and will tend to sink. By contrast, the mantle lithosphere portion must be less dense than the ambient mantle (PREM) because it contains: (1) a thick section (< 30 to 60 km) of MORB melting residues beneath the crust that are less dense, (2) water because of incomplete subduction-dehydration (Niu, 2004), which is hosted in serpentinite before subduction (developed near ridges and the trench-outter rise), in DHMS phases during subduction, in perovskite in the lower mantle, in ringwoodite/wadsleyite in the Transition Zone, and in olivine in the upper mantle (or a vapour phase, causing incipient melting), which reduces both the bulk-rock density and the elastic moduli (e.g., Frost, 1999; Litasov et al., 2003; Mao et al., 2008a,b; Ye et al., 2009; Li et al., 2009; Jacobsen et al., 2008), and (3) refertilized volatile-rich deep portions of the oceanic lithosphere (Figure 7). This means that the subducted lithosphere slab will likely separate into the dense crust (SOC) that sinks to the base of the mantle to form the LLSVPs and the buoyant mantle lithosphere of the slab (SML) that remains at shallower mantle depths. It is thus possible that the dense LLSVPs (SOC) could be overlain by such buoyant subducted mantle lithosphere (SML; Figure 10). Because the SML is ~ 15 times the mass of the SOC, it can occupy much of the lower and perhaps part of the upper mantle, and because it is buoyant relative to the ambient mantle (e.g., PREM), it will result in a surface manifestation – the geoid high and superswell in the Pacific and the “poorly-understood high elevation” of the African continent.

Why do LIP eruption sites correspond to edges of the LLSVPs?

Burke and Torsvik (2004) found that 24 active hotspot volcanoes can be projected to the DVs = -1% contours along the edges of the LLSVPs (also see Thorne et al., 2004; Garnero et al., 2007). Burke et al. (2008) further showed that all LIP (large igneous province) eruption sites with ages < 300 Ma lie above the DVs = -1% contours at the LLSVP edges, which they called plume generation zones (PGZs), suggesting that the LLSVPs may have remained unchanged for at least the past 300 Ma. The latter is expected if the LLSVPs are indeed piles of SOC because (1) ocean crust of MORB composition is much denser than the ambient materials at the base of the mantle in both solid state and liquid form (see Figure 9; also Figure 7 of Niu and O’Hara, 2003), and (2) if our calculations are reasonable, then at 300 Ma, the LLSVPs would already have ~ 92.5% of their present mass/volume. Localized ultra-low velocity zones (ULVZs) beneath, or in the vicinity of, the LLSVPs just above the CMB (e.g., Garnero et al., 1998; Williams et al., 1998) may be localized melt layers within the SOC induced by core heating. Such melt would be too dense to rise (Figure 9).

It is puzzling why LIP eruption sites should correspond to the edges (vs. centres/interiors) of the LLSVPs. Burke et al. (2008) speculated that “hot material that has been heated by conduction from the core in the basal part of the slab graveyard may be driven toward the PGZs (i.e., the edges of the LLSVPs) by slabs or slab fragments acting like push brooms.” Our explanation is much simpler as shown in Figure 10. At the edge of the LLSVPs, which act as thermal insulators, the subducted mantle part of the lithosphere (SML) loses its negative buoyancy by heating from the core and rises. Water may facilitate partial melting in this hot setting, and because peridotite melt is ~ 12% less dense than the ambient mantle (e.g., PREM, see Figure 9), this partially molten peridotitic system will readily rise as diapirs (or initial “plumes” if one prefers to call so), explaining why major hotspot volcanoes and LIP eruption sites correspond to the edges (vs. the interiors) of the LLSVPs. Some “small” volcanoes do occur on the topographic highs or swells, for which our interpretation is given in Figure 10 (represented by “g”).

If the above diapirs are indeed the source for OIB, then, as we have argued above, the OIB source materials are likely peridotitic in composition and could include the recycled deep portions of ancient
3 Subducted continental sediments (CC) can be introduced into source regions of OIB, but there is no convincing evidence for their significance as a major enriched component in the global OIB source in terms of the familiar CC-like signatures. One possibility is that subducted continental sediments as well as subduction-eroded continental materials may have been largely mixed with and diluted by the ambient mantle with the otherwise distinct CC-like signatures lost before transported to OIB source regions.

4 The observation that OIB are not only enriched in incompatible elements, but also enriched in the progressively more incompatible elements with respect to major

Summary

1 The geochemically enriched trace element signatures of OIB are largely inherited from their enriched fertile mantle source materials, including recycled deep portions of subducted oceanic lithosphere (“T” in Figure 7), but can also be enhanced by mixing with a melt-rich layer atop the LVZ (“II” in Figure 7) and by assimilation of earlier-formed metasomatic vein lithologies (“III” in Figure 7) in the oceanic lithosphere prior to eruption. A peridotite source (“T”) is required, and (“II”) and (“III”) are not; however, the participation of the latter two can lead to extremely enriched OIB compositions (e.g., basanite and nephelinite melt).

2 Subducted ocean crust (SOC) is too depleted (i.e., [La/Sm]_N < 1) to act as a major source material for highly enriched OIB (e.g., [La/Sm]_N >> 1) (Figs. 1-3), which is not surprising because SOC, once entering the lower mantle, is too dense to rise back again to upper mantle depths (Figure 9). The revised ROC-model involving olivine-free pyroxenite as the source for Hawaiian OIB is apparently attractive, but has many problems. For example, this olivine-free pyroxenite, postulated to result from the interaction of SOC and harzburgite remains too depleted to explain the highly enriched geochemical signature of OIB.

3 Subducted continental sediments (CC) can be introduced into source regions of OIB, but there is no convincing evidence for their significance as a major enriched component in the global OIB source in terms of the familiar CC-like signatures. One possibility is that subducted continental sediments as well as subduction-eroded continental materials may have been largely mixed with and diluted by the ambient mantle with the otherwise distinct CC-like signatures lost before transported to OIB source regions.

The observation that OIB are not only enriched in incompatible elements, but also enriched in the progressively more incompatible elements with respect to major

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mantle minerals, and that OIB sources are, in general, more enriched than the primitive mantle requires that OIB source materials be pre-enriched by low-F melt metasomatism. With all the possibilities considered, the interface (i.e., LAB) between the growing oceanic lithosphere and the top of the LVZ is the ideal site for mantle metasomatism, generating geochemically enriched deep portions of oceanic lithosphere. Globally, the 70 Myr history of oceanic lithosphere accretion to its full thickness of 90 km from the underlying LVZ records the history of mantle metasomatism. Similar mantle metasomatism may also take place at the base of the continental lithosphere. It is possible that partial melting may take place in the mantle transition zone, especially in the vicinity of the 410 km seismic discontinuity, but whether such melt, if formed at all, possesses the enriched geochemical characteristics as a metasomatic agent for OIB requires investigations.

5 The LVZ is compositionally layered. The reduction in seismic velocity at, or immediately below, the lithosphere-LVZ interface (i.e., LAB) is attributed to an H2O-CO2-rich silicate melt derived from within the LVZ. This melt is enriched in incompatible elements and migrates upwards, concentrating atop the LVZ. The deep portion of the LVZ thus becomes progressively more depleted and serves as the source for MORB.

6 In the subducted oceanic lithosphere, the SOC is very dense and will sink, contributing to the development of the LLSVPs at the base of the mantle, most prominently beneath the Pacific and Africa. The buoyant (relative to PREM) mantle part of the lithosphere (SML), which is ~ 15 times the mass of the SOC, tends to “float”, overlying the two global LLSVPs, explaining the surface geoid highs of the Pacific superswell and the elevated African continent and the associated dispersed volcanism.

7 The LLSVPs act as thermal insulators, which prevent the overlying SML from being heated and melted. However, at the edges of the LLSVPs, the SML material can be heated more directly by the outer core, and can melt. The low density of the SML (vs. PREM) and the even lower density of peridotite melts can cause the SML at the edges of the LLSVPs to form compositionally buoyant diapirs. These diapirs may become volumetrically significant and when they rise into the upper mantle, they will become significant melt sources, explaining why the eruption sites of major hotspot volcanoes and LIPs are associated with the edges, not interiors, of the LLSVPs.

8 We emphasize again that the major fertile OIB source material from depth (i.e., “T” in Figure 7) is likely SML, in particular the deep portions of metasomatic origin (Figure 7) because of their relative buoyancy and enriched geochemical characteristics; its bulk composition is peridotitic. On the other hand, subduction of the ocean crust is irreversible because of its high density; hence, there is little recycled SOC involved in the OIB petrogenesis (although tiny fragments of SOC could be entrained in some form of upwellings).

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