Paleo-Asian oceanic slab under the North China craton revealed by carbonatites derived from subducted limestones

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ABSTRACT

It is widely accepted that the lithospheric mantle under the North China craton (NCC) has undergone comprehensive refertilization due to input from surrounding subducted slabs. However, the possible contribution from the Paleo-Asian oceanic slab to the north is poorly constrained, largely because of the lack of convincing evidence for the existence of this slab under the NCC. We report here carbonatite intruding Neogene alkali basalts in the Hannuoba region, close to the northern margin of the NCC. Trace element patterns with positive Sr and U anomalies, negative high field strength elements (Nb, Ta, Zr, Hf, and Ti) and Ce anomalies, high \(^{87}\text{Sr}/^{86}\text{Sr}\) ratios (0.70522–0.70796), and high \(^{8}\text{O}_{\text{SMOW}}\) (standard mean ocean water) values (22.2‰–23‰) indicate that this carbonatite had a limestone precursor. However, the presence of coarse-grained mantle-derived clinopyroxene, orthopyroxene, and olivine, and chemical features of the carbonates suggest that the carbonate melts were derived from the mantle. The carbonates have high \(^{143}\text{Nd}/^{144}\text{Nd}\) ratios (0.51282–0.51298) and show negative correlation between \(\text{CaO}\) and \(\text{Ni}\) contents, resulting from reaction between carbonate melt and peridotite. Considering the regional tectonic setting, the carbonatite probably formed by melting of subducted sedimentary carbonate rocks that formed part of the Paleo-Asian oceanic slab, and thus could provide the first direct evidence for the presence of the Paleo-Asian oceanic slab beneath the NCC.

INTRODUCTION

Carbonate platforms are common in oceanic realms. Small tropical oceans, exemplified by many segments of the Tethys Ocean in Europe, were covered with extensive carbonate platforms, found today in classic Alpine fold and thrust belts. It is clear that some of these materials are subducted to mantle depths where they undergo decarbonation reactions and may be involved in partial melting (Collins et al., 2015; Doucelance et al., 2014; Ducea et al., 2005; Hammouda, 2003). Enormous budgets of CO\(_2\) at magmatic arcs (Lee and Lackey, 2015) independently require that carbonate is very influential in mass exchange at convergent margins. Subduction magmatism is probably one of the most important tectonic mechanisms responsible for regulating the exchange of CO\(_2\) between the Earth’s interior and the atmosphere (McKenzie et al., 2016). Unfortunately, very few observational data exist to provide details on the mechanisms of transport of carbonate materials and CO\(_2\) from the surface to mantle depths and back to the Earth’s surface via fluid (decarbonation reactions) and melt transport (Liu et al., 2015).

It has been suggested that the lithosphere of the North China craton (NCC) was successively modified by (1) the southward subduction of the Paleo-Asian oceanic slab (PAOS) between the early Paleozoic and the late Permian, (2) the northward subduction of the Paleo-Tethys oceanic and Yangtze plates in the Triassic, and (3) the westward subduction of the Pacific plate since the Cretaceous (Fig. 1A) (Windley et al., 2010). Although all of these subduction events could have contributed to widespread lithospheric reactivation and thinning under the NCC (e.g., Griffin et al., 1998), the Pacific plate subduction was generally advocated to account for the lithospheric thinning (Zhu et al., 2011). The contribution of the PAOS and Paleo-Tethys oceanic slab to lithospheric thinning remains enigmatic, largely because of the lack of convincing evidence for the existence of these slabs under the NCC. If abundant carbonate sediments were transported into deep mantle during oceanic slab subduction, they could have contributed fundamentally to the modification of the chemical and physical properties of the lithospheric mantle by carbonate metasomatism. In turn, subsequent mantle-derived melts (especially carbonatite) could contain the fingerprint of subducted carbonate, which can be used to trace the origin of the subducted slab (Tappert et al., 2009) and outline the crust-mantle recycling of carbonate sediments.

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Figure 1. A: Tectonic framework of the study area (modified from Windley et al., 2010). CAOB—Central Asian orogenic belt; QDSOB—Qinling-Dabie-Sulu orogenic belt. B: Field appearance of the carbonatite intrusion. C: Residual olivine (Ol) xenocryst partly resorbed by carbonatite melt. SC—sparry calcite; CM—carbonate matrix. D: Monomineralic aggregates consist of interlocking calcite grains.
Here we document a carbonatite intrusion with geochemical features of recycled limestone. The intrusion marks the subduction of an overlying carbonate platform of the PAOS, to mantle depths beneath the NCC, providing evidence for recycling of carbonate back to the Earth’s surface by buoyant diapirism and high-degree melting.

SAMPLES, PETROLOGY, AND GEOCHEMICAL COMPOSITIONS

Samples were collected from a carbonatite intrusion (0.5–3 m thick and >35 m wide) that intrudes Neogene basalts (22–10 Ma; Zhu, 1998) at Hannuoba (eastern China; Fig. 1). The basalts are distributed along the northern margin of the NCC, and comprise intercalated tholeiitic, transitional, and alkaline basalts (Zhi et al., 1990). The source of the basalts probably contains abundant garnet-pyroxenite (Liu et al., 2008) or meta-somatized by carbonate-rich fluid (Dupuy et al., 1992). The alkaline basalts carry abundant granulite, pyroxenite, and peridotite xenoliths (Chen et al., 2001). The evolved Sr-Nd isotopic compositions of some pyroxenite xenoliths indicate involvement of subducted sediments (e.g., Xu, 2002).

The carbonatite intrusion occurs in xenolith-free basalt and has a sharp boundary with the basalt layer. The overlying basalt is domed upward by the invasion of the carbonatite melt (Fig. 1B).

The carbonatite contains aggregates of calcite phenocrysts and medium- to coarse-grained silicate macrocrysts set in a matrix of fine-grained calcite (Figs. 1C and 1D), indicating rapid quenching. Monomineralic aggregates of interlocking calcite grains (50–100 μm) indicate calcite accumulation before quenching. The silicate macrocrysts consist of coarse-grained clinopyroxene (Cpx, 0–15 vol%), orthopyroxene (Opx, 0–12 vol%), olivine (Ol, 0–10 vol%), and rare spinel (Sp, 0–1.5 vol%). Ol is forsteritic (Fo90–91) and has low CaO contents (<0.1 wt% and K2O <0.05 wt%); many are similar to limestones, and some are along a mixing trend between limestone and peridotite (Fig. DR2). MgO contents in the carbonatites are higher than in limestones (0.13–1.65 wt%), and show a negative correlation with CaO. Trace element patterns are similar to sedimentary limestones with notably positive Sr and U anomalies and negative high field strength element (HFSE; Zr, Hf, Nb, Ta, Ti) anomalies (Fig. 2). The incompatible trace element contents of most samples are lower than average limestone, especially for Rb, Ba, and heavy rare earth elements (REEs) (Fig. 2). Their REE patterns show negative Ce anomalies (Ce/Ce* = 0.2–0.8) and positive Eu anomalies (Eu/Eu* = 1.15–2.98) (Table DR2). The carbonatite matrix contains 45.2–52.7 wt% CaO, 0.4–1.8 wt% MgO, and 1.5–5.0 wt% SiO2 (Table DR3). Both the carbonate matrix and phenocrysts have higher Ni contents (1–133 ppm) than limestone, and show negative correlations between CaO and Ni (Fig. 3A).

Carbonates in the carbonatite intrusion have higher 87Sr/86Sr (0.70522–0.70796) and slightly higher 143Nd/144Nd (0.51282–0.51298) than typical carbonatites (Fig. 3B). 143Nd/144Nd ratios show no correlation with Nd contents (0.4–2.8 ppm), whereas 87Sr/86Sr ratios correlate negatively with Sr contents (27–306 ppm) (Fig. DR3). Rare aragonite veinlets that crosscut one sample have high Sr contents (2507–6600 ppm) and relatively low 87Sr/86Sr ratios (0.70411–0.70683) (Table DR4); the postmagmatic fluid that deposited this aragonite might have also partially modified the sample (Fig. 3B). The primary carbonatitic melts represented by the low-Sr samples have the highest 87Sr/86Sr and 143Nd/144Nd ratios. The carbonatites are characterized by light carbon isotopic compositions [δ13CVPDB (Vienna Peedee belemnite) = −14.4% to −11.2%] but heavy oxygen isotopic compositions [δ18O (SMOW—standard mean ocean water) = 22.2‰ to 23‰] (Table DR2).

DISCUSSION

Mantle Derivation of the Carbonatite

Possible origins of the Ol and Cpx macrocrysts in the carbonatite intrusion are reaction products between carbonate melt and the basalt wall rock (Jolis et al., 2013) or peridotite xenocrysts from the mantle that potentially provide records of the ascent process of the carbonatite melts. The compositions of Ol (CaO <0.1 wt%) and Cpx (Mg# ≈ 91.9–93.3 and SiO2 >50.1 wt%) macrocrysts clearly correspond to minerals in mantle peridotites and are unlike products
of carbonate melt–basalt interaction (Fig. DR4). This conclusion is supported by the Ni contents of Opx and Cpx (Table DR1), which are also typical for mantle minerals (Foley et al., 2006). OI produced by carbonate melt–basalt interaction would have higher CaO (>1.31 wt%), and Cpx would be characterized by low Mg# (<84) and low SiO₂ content (<45 wt%) (Jolis et al., 2013). The temperatures estimated from Cpx + Opx macrocryst pairs are 770–932 °C (see the Data Repository), implying a source depth of ~50 km for the disrupted mantle xenoliths (see the Data Repository). Taken together, these features suggest that the subducted carbonate precursor could have been partially modified by aqueous fluids in the subduction zone. Furthermore, the much lower heavy (H) REE contents and positive Eu anomalies in the carbonatites compared to average limestone (Fig. 2) might be attributed to modification by subduction-related CO₂-rich aqueous fluids. HREE” are complexed more strongly by CO₂ than LREE”, and similarly LREE” are complexed more strongly by CO₂ than Eu” (Bau, 1991), so that HREE” would be removed easily from limestone by a CO₂-rich fluid, but Eu” could be retained.

Melting of limestone is commonly regarded to be restricted to unusually hot regimes (Wylie and Tuttle, 1960), although the temperatures involved may be reduced in water-bearing conditions (Poli, 2015). Due to the high solidus temperatures but lower density and viscosity, limestone cannot be melted at the top of the subducted slab, but may penetrate the cold cratonic mantle in the form of solid buoyant diapirs (Behn et al., 2011). The large scale of basalt in the Hannuoba area (~1700 km²) indicates a giant Cenozoic thermal event in the mantle under the northern margin of the NCC that could have resulted in substantial melting of such limestone diapirs. Limestone is mostly composed of a single calcite phase, so it is reasonable to speculate that high modes of melting of subducted limestone would occur once melting initiates. High-degree melting of a subducted limestone in the mantle was also suggested (in Liu et al., 2015) to account for the mantle-derived carbonatite xenoliths from the Dalihu area with the geochemical features of limestone. The Hannuoba carbonatite intrusion may share an origin similar to that of the carbonatite xenoliths reported in Liu et al. (2015), but the intrusion was emplaced into the shallow crust. The carbonate melt might have undergone complex devolatilization during ascent (Russell et al., 2012), which could have produced the low δ¹³CVPDB values of carbonatite melt (Cartigny et al., 1998) and also yielded extremely local, strongly reducing conditions that permitted the formation of moissanite (Shiryaev and Gaillard, 2014). This would explain both the much lower δ¹³CVPDB values of the Hannuoba carbonatites relative to average limestone (−0.44‰ ± 2‰; 1σ, n = 102), and the occurrence of moissanites in the Hannuoba intrusion (Fig. DR1).

**Implications for the Subduction of the Paleo-Asian Oceanic Slab Under the NCC**

The δ⁸⁷Sr/⁶⁰Sr ratios of sedimentary carbonate rocks have varied with time (Veizer et al., 1999) (Fig. DR5), so that the high Sr isotopic compositions can be used to constrain the age of the limestone. δ⁸⁷Sr/⁶⁰Sr ratios of marine carbonates increased from ~0.702 in the Archean to ~0.7069 in the late Neoproterozoic, and varied within a small range (0.7069–0.7089) during the Phanerozoic. The highest δ⁸⁷Sr/⁶⁰Sr ratio (0.70791) of carbonate in the Hannuoba carbonatite represents materials least contaminated by aragonite veins and mantle materials (Fig. 3B), from which we infer that the age of the limestone source of the carbonatite is most probably between 580 and 360 Ma, and certainly not older than 580 Ma (Fig. DR5); this is consistent with its derivation from the Paleo-Asian Ocean, which existed between 1 Ga and 250 Ma (Xiao et al., 2003). Furthermore, the carbonatite intrusion occurs ~1000 km from the trench position of the east Pacific plate and the Qinling-Dabie-Sulu orogenic belt, but within 200 km of the northern margin of the NCC (Fig. 1A). The Pacific slab is eliminated as a possible source because it is stagnating in the mantle transition zone beneath the eastern NCC (Zhao et al., 2009) at a depth at which limestone could not be subducted due to its low density and viscosity. These observations suggest that the limestone precursor was most probably derived from the subducted PAOS.

It is noteworthy that Mesozoic carbonatites from Zhulu and Huairen ~50–100 km south of Hannuoba have high δ⁸⁷Sr/⁶⁰Sr ratios (0.7055–0.7075; Yan et al., 2007) (Figs. 1 and 3B) similar to those of the Hannuoba carbonatite intrusion, implying that they may also be related to the PAOS. Although the Mesozoic Zhulu and Huairen carbonatites have similarly lower trace element contents than typical carbonatite, they have much higher trace element contents than the Cenozoic Hannuoba intrusion (Fig. 2). This suggests that the Mesozoic carbonatites could have been derived from partial melting of subducted carbonate-rich eclogite rather than limestone. The Hannuoba, Zhulu, and Huairen carbonatites are distributed in a north-south direction, and show increasing trace element contents with increasing distance from the north margin of the NCC (Figs. 2 and 4). The temporal and spatial variations of the Mesozoic and Cenozoic carbonatites are consistent with the interpretation that carbonatitic eclogite could have been subducted deep into the mantle (Hammouda, 2003) before melting, whereas limestone would have detached...
from the downgoing slab to form buoyant diapirs penetrating the shallow and cold mantle (Behn et al., 2011; Liu et al., 2015) (Fig. 4). Subsequently, a giant Cenozoic mantle thermal event, as indicated by the large-scale Hannuoba basalts, triggered high-degree melting of the limestones, now in the shallower mantle. These carbonatites reconstruct the path of the consumed PAOS under the NCC, which is critical for understanding how subduction zones modulate the global carbon cycle.

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