Special Topic: Tracing Deep Carbon Cycles by Metal Stable Isotopes

Linking deep CO₂ outgassing to cratonic destruction

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ABSTRACT

Outgassing of carbon dioxide from the Earth’s interior regulates the surface climate through deep time. Here we examine the role of cratonic destruction in mantle CO₂ outgassing via collating and presenting new data for Paleozoic kimberlites, Mesozoic basaltic rocks and their mantle xenoliths from the eastern North China Craton (NCC), which underwent extensive destruction in the early Cretaceous. High Ca/Al and low Ti/Eu and δ²⁶Mg are widely observed in lamprophyres and mantle xenoliths, which demonstrates that the cratonic lithospheric mantle (CLM) was pervasively metasomatized by recycled carbonates. Raman analysis of bubble-bearing melt inclusions shows that redox melting of the C-rich CLM produced carbonated silicate melts with high CO₂ content. The enormous quantities of CO₂ in these magmas, together with substantial CO₂ degassing from the carbonated melt–CLM reaction and crustal heating, indicate that destruction of the eastern NCC resulted in rapid and extensive mantle CO₂ emission, which partly contributed to the early Cretaceous greenhouse climate episode.

Keywords: carbonate metasomatism, lithospheric mantle, CO₂ outgassing, deep carbon cycling, cratonic destruction, North China Craton

INTRODUCTION

Carbon exchange between the Earth’s interior and exterior exerts an important influence on the surface climate through geologic time and is critical for planetary habitability. In recent years, it has been increasingly recognized that the cratonic lithospheric mantle (CLM) stores vast amounts of carbon, resulting from gradual enrichment by upward melt infiltration, in addition to the original carbon incorporated during its formation [1,2]. Carbon in the CLM can be extensively remobilized and released via continental rifting [1,3], active island arc volcanism [4,5] and plume-related magmatism [6,7], which represent three main ways proposed for mantle CO₂ emission. For example, up to 28 to 34 Mt of carbon per year (expressed as Mt C yr⁻¹) may be released by continental rifting [1]. A quantitative flux estimate for the CO₂ outgassing along with the massive Tan-Lu Fault Belt in eastern China gave 70 ± 58 Mt C yr⁻¹ [8]. Extensive CO₂ degassing (71 ± 33 Mt C yr⁻¹) has been estimated through extensional faults along the entire East African Rift [3], which is even comparable to the estimates for CO₂ degassing in island arcs (18–43 Mt C yr⁻¹) and mid-ocean ridges (8–42 Mt C yr⁻¹) [9–11]. Plume-induced CO₂ outgassing has also been proposed to have the ability to have caused abrupt climate changes in Earth’s history [12].

Cratons commonly retain tectonic and magmatic quiescence for billions of years [13], but some cratonic regions record extensive crustal deformation and on-craton magmatism that reflect cratonic destruction processes [14,15]. In sharp contrast to many others on Earth, the eastern part of the North China Craton (NCC) is a reactivated craton with the present-day lithosphere being made up of decoupled crust and mantle, i.e. Archean-Proterozoic crust and Phanerozoic lithospheric mantle [15]. The Archean thick (diamond-bearing) and cold lithospheric keel (>200 km) was partially or even wholly destroyed and removed, and was then replaced by a newly formed thin and hot lithospheric mantle (∼75 km), resulting in up to ∼120 km of the lithospheric keel being lost [14,16]. Reactivation of the CLM gave rise to a magmatic peak at ∼125 Ma including both mafic and felsic magmatism,
marking the climax of cratonic lithospheric destruction in the early Cretaceous [16,17]. Since destruction/thinning is confined to the eastern part of the NCC (its western part remains largely intact), the westward subduction of the paleo-Pacific oceanic slab underneath the eastern Asian continent in the early Cretaceous was widely advocated to have resulted in reactivation and destruction of the eastern NCC [18].

In order to examine whether cratonic lithospheric destruction results in massive mantle CO2 outgassing, we firstly ascertain whether or not the CLM beneath the eastern NCC was initially carbon rich and whether it had been widely subjected to carbonate metasomatism, including carbon from recycled carbonates prior to destruction in the early Cretaceous. For this purpose, we analyzed magnesium (Mg) isotopes for early Cretaceous lamprophyres and collated available chemical and Mg isotopic data for mantle xenoliths in Paleozoic diamond-bearing kimberlites and early Cretaceous mafic igneous rocks as well as orogenic ultramafic massifs in the Dabie orogen located on the south margin of the eastern NCC (Fig. 1). Then, we analyzed the CO2 components of melt inclusions (MIs) in early Cretaceous lamprophyres that can be used to calculate the CO2 concentrations in pre-eruptive magmas. Finally, we considered carbonated silicate melt–CLM reaction and crustal heating as additional ways for mantle CO2 outgassing to occur during cratonic destruction, in addition to mafic magmatism. Our results show that the CLM beneath the eastern NCC had widely interacted with carbonated melts prior to the early Cretaceous and
extensive CO₂ emission had occurred as a consequence of cratonic destruction.

**PRIMORDIAL CARBON IN THE CLM**

The terrestrial mantle initially contained carbon resulting from accretion and core-mantle differentiation processes [11]. Recent studies provide a rigorous reconstruction of carbon concentration for the MORB source mantle and suggest that the upper mantle contains ∼30 ppm C [19]. In addition to the primordial carbon, the continental lithospheric mantle, mainly formed between 2 and 3 Ga, contains more carbon (∼89 ppm) that is incorporated into the sub-arc lithosphere via the accretion of island arcs [1]. Assuming an area of ∼1000000 km² and lithospheric mantle thickness of ∼150 km for the eastern NCC prior to thinning in the Mesozoic, the CLM beneath the eastern NCC initially contains 4.27 × 10⁷ Mt C. In addition, the model of Foley and Fischer [1] predicts a long-term (>2 Ga) solid storage of carbon in the CLM as a result of episodic melt infiltration and redox freezing. If one considers this gradual enrichment from episodic freezing throughout the long evolution history of the ancient NCC (as old as ∼3.8 Ga; [20]), carbon concentration in the CLM beneath the eastern NCC may become much higher. Abundant diamonds were discovered in Paleozoic kimberlites from the NCC [21] and are direct evidence for a C-bearing, reduced CLM beneath the eastern NCC prior to the Mesozoic era. For example, the eruption of diamond-bearing kimberlites and the high Mg° (>90; 100 × mol C/ (Mg + Fe²⁺)) of olivines in diamond inclusions and xenolith/xenocryst olivines from Mengyin and Fuxian in the eastern NCC (Fig. 1) indicate the existence of a thick, low-density, cold root, which is mainly composed of refractory harzburgite and lherzolite [21].

**ADDITIONAL CARBON FROM RECYCLED CARBONATES**

Since the Paleozoic, the NCC further underwent multiple oceanic plate subductions from the south, north and east sides, which potentially added surface carbon into the CLM beneath it. Below we present lines of evidence for more recent carbon addition to the CLM beneath the eastern NCC, from recycled carbonates related to slab subduction.

**Finding of low-δ²⁶Mg lamprophyres**

Magnesium isotopes are a novel and efficient tool for identifying recycled carbonates that are isotopically much lighter than the mantle [22,23]. Lamprophyres are typically characterized by a high content of volatiles and commonly record fluid/melt–mantle interaction in their magma sources [24], thereby providing an opportunity to investigate the nature of fluids/melts responsible for CLM metasomatism. Early Cretaceous lamprophyres are widely exposed in the NCC and have a magmatic peak at ∼125 Ma (Fig. 1), which is contemporaneous with the climax of cratonic lithospheric destruction of the eastern NCC [25]. Here we present a Mg isotopic dataset for Shandong lamprophyres, and for comparison we collate chemical and Sr-Nd isotopic data for other early Cretaceous lamprophyres widely distributed in the eastern NCC (Fig. 1; Tables S1–S3). Some High-Ti lamprophyres from Shandong have relatively depleted Sr and Nd isotopic compositions (Fig. 2a) and were proposed to have been derived from the asthenospheric mantle [26]. Most of the early Cretaceous lamprophyres in the eastern NCC, including those reported in this study, however, have extremely enriched Sr and Nd isotopic compositions (87Sr/86Sr(t) = 0.70520–0.71099, εNd(t) = −18.8 to −8.3) that are in sharp contrast to the High-Ti lamprophyres and Cenozoic alkali basalts in the NCC (Fig. 2a), pointing to an enriched CLM source. According to MgO contents, the Shandong lamprophyres are classified into Low-MgO (MgO < 7.5 wt%) and High-MgO (MgO > 7.5 wt%) subgroups (Fig. 2). Low-MgO lamprophyres have mantle-like δ²⁶Mg (−0.32‰ to −0.24‰), whereas High-MgO lamprophyres possess significantly lower δ²⁶Mg (−0.59‰ to −0.35‰) than the mantle δ²⁶Mg value of −0.25 ± 0.04‰ (Fig. 2). It has been well demonstrated that Mg isotope fractionation during mantle partial melting and magma differentiation is limited (<0.07‰) [23]. In fact, the negative correlation between δ²⁶Mg and MgO (Fig. 2c) argues against the light δ²⁶Mg of the High-MgO lamprophyres being a result of fractional crystallization of any minerals involving removal of isotopically heavy Mg. More discussions about the influence of magma differentiation, as well as crustal contamination on Mg isotopic systematics of the studied lamprophyres, are provided in Supplementary Data. Overall, the variation of δ²⁶Mg in Shandong lamprophyres reflects isotopically heterogeneous mantle sources caused by recycled crustal carbonates.

A carbonated mantle source for High-MgO lamprophyres is corroborated by their systematically higher CaO (High-MgO, 9.47 ± 0.84 wt%; Low-MgO, 7.25 ± 1.37 wt%; 1sd; Fig. S1) and CaO/Al₂O₃ ratios (High-MgO, 0.69 ± 0.07; Low-MgO, 0.45 ± 0.1; Fig. 2d) and lower Ni
Figure 2. Plots of (a) $^{87}\text{Sr}/^{86}\text{Sr}_{i}$ versus $\epsilon_{\text{Nd}}(t)$ and (b) MgO versus CaO/Al$_2$O$_3$ for the lamprophyres, early Cretaceous volcanic rocks and Cenozoic basalts in the NCC. The gray circles represent the data of lamprophyres in previous studies, the gray triangles represent the data of Cenozoic basalts and the white diamonds represent the data of early Cretaceous volcanic rocks (all data are listed in Tables S1–S3). Plots of $\delta^{26}\text{Mg}$ versus (c) MgO, (d) CaO/Al$_2$O$_3$, (e) $^{87}\text{Sr}/^{86}\text{Sr}_{i}$ and (f) $^{143}\text{Nd}/^{144}\text{Nd}$ for the studied lamprophyres. Early Cretaceous basalts from Yixian, Feixian and Fangcheng (YX-FX-FX, yellow circles) in the eastern NCC [29] are shown for comparison. The magnesium isotopic composition of the terrestrial mantle (expressed as $\delta^{26}\text{Mg}$ in per mil relative to DSM3) is from ref. [23]. The results of Mg-Sr isotopic modeling show a two-stage source metasomatism: the first stage is associated with siliciclastic sediments, and the second stage is related to carbonate metasomatism. The parameters used for Mg-Sr isotopic modeling are listed in the Supplementary Data (Table S4).

Carbonates (e.g. dolomites; Fig. 2e). During slab subduction, Ca-rich carbonates can be substantially dissolved by aqueous fluids at initial stages and injected into the sub-arc mantle [9]. At larger depths of $>160$ km, dolomite dissolution occurs in subducting slabs and can be further enhanced by supercritical fluids [28]. The lithospheric mantle of the eastern NCC had a thickness of $>200$ km prior to thinning in the early Cretaceous [14,15]. Thus, the finding of low-$\delta^{26}\text{Mg}$ lamprophyres demonstrates that the CLM beneath the eastern NCC had been metasomatized by dissolved magnesium carbonates from subducting slabs. The Low-MgO rocks are SiO$_2$ rich and CaO poor and represent partial melts of the CLM metasomatized by recrystallized siliciclastic sediments, which explains their mantle-like $\delta^{26}\text{Mg}$ yet highly radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ compositions (Fig. 2). We suggest a two-stage source metasomatism: the first stage is associated with siliciclastic sediments that led to the enriched Sr and Nd isotopic signatures, and the second stage is related to carbonate metasomatism that injected the low $\delta^{26}\text{Mg}$ signatures, without significantly affecting Sr-Nd isotopic compositions (Fig. 2e and f). Previous studies found that early Cretaceous basalts from Fangcheng, Yixian and Feixian in the eastern NCC have mantle-like $\delta^{26}\text{Mg}$ [29] (Fig. 2c and d). It is noteworthy that these basalts have CaO/Al$_2$O$_3$ and CaO/TiO$_2$ ratios similar to those of the Low-MgO lamprophyres with normal $\delta^{26}\text{Mg}$. This indicates that the carbonated CLM may be chiefly sampled by lamprophyres that are commonly derived from a volatile-rich source [24]. Indeed, most of the Low-Ti lamprophyres with enriched Sr-Nd isotopic compositions from other regions in the eastern NCC have high CaO/Al$_2$O$_3$ and MgO content, which resembles the Shandong High-MgO lamprophyres with low $\delta^{26}\text{Mg}$ (Fig. 2a and b). This implies that the pre-Cenozoic CLM beneath the eastern NCC may have undergone widespread metasomatism by recycled carbonates. Carbonate minerals (e.g. magnesite and calcite) are often observed in coeval lamprophyres in the NCC [30]. Overall, our new Mg isotopic data provide solid evidence for a recycled carbonate component in the CLM beneath the eastern NCC at or prior to the early Cretaceous.

Evidence from xenoliths, ultramafic massifs and carbonatites

Mantle-derived xenoliths in volcanic rocks and orogenic ultramafic massifs sample the lithospheric mantle, serving as a direct window to observe mantle metasomatism. A large number of lherzolite, wehrlite and clinopyroxenite xenoliths carried by
Paleozoic diamond-bearing kimberlites, Mesozoic and Cenozoic mafic igneous rocks, and ultramafic massifs in the Dabie orogen derived from the deep mantle wedge (>160 km) on the south margin of the NCC (Fig. 1), record carbonate metasomatism of the CLM beneath the NCC. Wehlrites represent rocks where all, or most, orthopyroxene has been consumed through metasomatic reactions and are considered to be one of the end products of carbonate metasomatism in the CLM [8,31]. Pyroxenites and garnet pyroxenites represent rocks where all olivine and orthopyroxene have been consumed through metasomatic reactions with SiO₂ carried by supercritical fluid or silica-rich melt and are therefore considered to be other end products of carbonate metasomatism in the CLM. Abundant wehlrite xenoliths have been found in Mesozoic basaltic rocks from Tietonggou and Liaoyuan (Fig. 1a) [32,33], which suggests pervasive carbonate metasomatism of the CLM. Pyroxenite xenoliths hosted by the Jiagou intrusion (~130 Ma) in the southeastern NCC (Fig. 1) have extremely low δ²⁶Mg of −1.23‰ to −0.73‰ (Fig. 3a) [34]. These pyroxenite xenoliths have a metasomatic U-Pb isotopic age of ~400 Ma, suggesting carbonate metasomatism induced by paleo-Tethys slab subduction. Garnet pyroxenites in the Maowu ultramafic massif have low δ²⁶Mg of −0.99‰ to −0.65‰ and contain abundant carbonate mineral inclusions and metasomatized zircons with high δ¹⁸O (up to 12.2‰), suggesting metasomatism of the CLM by recycled carbonates. The age of zircons (457 ± 55 Ma) from the garnet clinopyroxenites also indicates Paleozoic metasomatism by subduction of the paleo-Tethys ocean slab [28]. Some pyroxenite and garnet pyroxenite xenoliths hosted by Cenozoic basalts (e.g. Hannuoba) also have low δ²⁶Mg [35] (Fig. 3a). Because Hannuoba is located to the west of the Daxing’anling-Taibai gravity lineament (DTGL) in the western part of the NCC, in which the CLM has not been affected by the Mesozoic thinning, the presence of low-δ²⁶Mg xenoliths also indicates mantle carbonate metasomatism of the NCC prior to Cenozoic. It is noted that the Hannuoba xenoliths have δ²⁶Mg (low to −1.42‰) (Fig. 3a) much lower than those of the host basalts and all other Cenozoic basalts in eastern China (−0.6 to −0.3‰) [29], thus their low δ²⁶Mg is unlikely to have been caused by interaction between low-δ²⁶Mg basaltic melt and the overlying lithospheric mantle. Generally, there is a negative correlation between δ²⁶Mg and CaO content for these xenoliths (Fig. 3b), strongly suggesting metasomatism of the CLM by recycled carbonates. Apart from Mg isotopes, Ca/Al, (La/Yb)N and Ti/Eu ratios of clinopyroxenes are effective indices of mantle carbonatitic metasomatism. As shown in Fig. 3c and d, clinopyroxenes in mantle xenoliths hosted by Paleozoic diamond-bearing kimberlites and Mesozoic mafic igneous rocks have systematically higher Ca/Al and (La/Yb)N and lower Ti/Eu ratios than those of silicate-metasomatic mantle xenoliths and deplet-MORB-mantle (DMM) peridotites. Along with high Mg and low Ti/Eu, these xenoliths are believed to have undergone carbonatitic metasomatism [36].

Further evidence for pre-Cenozoic carbonate metasomatism of the CLM comes from carbonatites. The solidus of mantle rocks can be reduced by addition of volatiles such as CO₂ into the mantle and melting of the CO₂-rich mantle would produce alkali-rich and silicon-poor melts, such as carbonatites [27]. Mesozoic carbonatites are exposed at more than 10 locations in the NCC [36–38] (Fig. 1). The enriched Sr and Nd isotopic compositions of these carbonatitic magmas suggest an enriched, carbonated mantle source [37]. Mesozoic carbonatites from Zhuolu and Huairen have high ⁸⁷Sr/⁸⁶Sr ratios (0.7055–0.7075) and are proposed to have formed by direct melting of recycled sedimentary carbonates in the mantle [38]. Carbonatites intruding on Neogene alkali basalts in Hannuoba on the northern margin of the NCC have high ⁸⁷Sr/⁸⁶Sr (0.70522–0.70796) and high ¹⁸O ratios (22.2‰–23.0‰), which are directly linked to the subducted...
Recycled carbonates would be reduced to diamond or graphite at depths of >150 km [40].

Collectively, the lines of evidence above strongly suggest that the CLM beneath the eastern NCC has been subject to pervasive carbonate metasomatism since the Paleozoic. The carbonate metasomatism could have been induced by multiple oceanic plate subduction events around the NCC, that is, the paleo-Asian oceanic slab in the north, paleo-Tethys oceanic slab in the south and paleo-Pacific slab in the east (Fig. 4). These subducted slabs carried large amounts of carbonate sediments into the mantle and transformed the CLM into a vast store for carbon. However, the deep part of the mantle is commonly too reduced to favor stable carbonates. That is, when carbonates are recycled into the mantle at depths of >120 km, they will be reduced via the following redox reaction:

\[
\text{MgSiO}_3 + \text{MgCO}_3 = \text{Mg}_2\text{SiO}_4 + \text{C} + \text{O}_2.
\]

At depths of 120–170 km, recycled carbonate is transformed into carbon that exists as graphite and at larger depths (>170 km) as diamond [40], although in the CLM diamond is stable to lower pressures at cool conductive geotherms (Fig. 4). It is difficult to quantify the flux of recycled carbon in the mantle of the entire NCC since the Paleozoic, but we can give a rough estimate for this study area. As discussed above, the Mg-Sr isotopic mixing model indicates that the mantle source of low-δ^{26}Mg lamprophyres contains ~10 wt% Mg-rich carbonates (Fig. 2e), which is roughly equivalent to ~1 wt% C. Assuming a density of 3.2 g cm⁻³ and a possible 40-km lithosphere depth interval that has been metasomatized, the mass of recycled C in the CLM beneath Shandong peninsula can be calculated. The lithosphere beneath the eastern NCC was >200 km thick before destruction [18,41] and the depth at which Mg-rich carbonates start to dissolve is ~160 km [28,29]; thus we assume an ~40-km interval for carbonate metasomatism. An areal estimate is available for the Shandong peninsula (∼73 000 km²), and we assume that about half the area was affected based on the proportion of occurrence of High-MgO lamprophyres with low δ^{26}Mg in the study area (Fig. 2). From this, 6.09 × 10⁷ Mt C is estimated to have been added by recycled carbonates. Together with the primordial carbon (∼4.27 × 10⁷ Mt C) in the CLM prior to the Paleozoic, the total reservoir of carbon in the CLM beneath the eastern NCC would be 1.04 × 10⁸ Mt C at least, which represents a significant store of carbon in the CLM with important contribution from recycled carbonates. The reduced CLM, with carbon mainly existing as graphite or diamond, has not undergone redox melting and was preserved until the Mesozoic during which it was largely activated and removed.

**DEEP CO₂ OUTGASSING INDUCED BY CRATONIC DESTRUCTION**

Commonly, the deep CLM is primarily reduced as a result of depletion in basaltic melt and the pressure effect on the oxygen fugacity during its formation [42]. A reduced CLM beneath the thick NCC (>200 km) is indicated by the Paleozoic diamond-bearing kimberlites. However, it can become oxidized as the diamondiferous CLM is exhumed to shallower depths due to lithospheric thinning, extension and mantle upwelling [8]. During this process, ‘redox melting’ would occur and carbon (diamond or graphite) in the CLM would become unstable and be oxidized by the reduction of Fe^{3+} at depths of <170 km [40]. This redox melting would produce carbonatitic melts at depths of ~150 km that could evolve into carbonated silicate melts accompanied by silicate melting at shallower depths. The redox melting of the CLM was probably induced by the decompression and rise of the lithosphere–asthenosphere boundary due to slab rollback of the westward subducting paleo-Pacific plate at the early Cretaceous [18]. Carbonatitic melts have much lower viscosity and density relative to silicate melts [43], which could further promote carbonate metasomatism of the shallower CLM. In the presence of carbonatitic melts, the mantle could be readily fusible, leading to efficient extraction of carbon from the deep interior [11]. Therefore, given the extensive thinning (>120 km) of the lithospheric mantle keel of the eastern NCC...
[14,16], extensive CO₂ outgassing is expected to have occurred as the CLM underwent redox melting during this thinning process.

Here we evaluate whether or not the early Cretaceous lamprophyres are CO₂ rich by analyzing gas exsolution bubble-bearing MIs in them. MIs are small droplets of silicate melt trapped by crystals in magmatic rocks and can be used to constrain the contents of volatile components dissolved in melts prior to volcanic eruption and, ideally, degassing. After the MIs are captured, bubbles will be formed during the cooling process of melts, post-entrapment crystallization on MI walls, or diffusive H⁺ loss [44]. Thus, the CO₂ of MIs is present mainly in bubbles due to its low solubility in silicate melts if post-entrapment degassing occurs [44]. MIs are mainly hosted in clinopyroxene and occasionally in olivine and amphibole macrocrysts of the Shandong lamprophyres (Fig. S3). The compositions of gas exsolution bubbles were analyzed by Raman spectroscopy (see Supplementary Data for detailed methods). Among the ∼200 MIs we analyzed, >80% of the MIs contain vapor bubbles and ∼20% of the vapor bubbles contain CO₂. The analyzed bubbles in most MIs are composed of pure or nearly pure CO₂ without other volcanic gases (CO, CH₄, H₂S, H₂O) being detected. The presence of CO₂ in the bubbles of MIs was confirmed by two characteristic peaks, at ∼1285 cm⁻¹ and ∼1388 cm⁻¹, defining a Fermi diad in the Raman spectrum (Fig. 5). CO₂ density (d) of the bubbles can be calculated by the spacing of the Fermi diad (Δ cm⁻¹), using the equation of Kawakami et al. [45]. The mass of CO₂ in bubbles can be calculated by multiplying CO₂ density by the volume of the bubble (Table S6). Then, the CO₂ content of the vapor bubble in ppm, [CO₂]vb, can be calculated using the following equation [44]:

\[
[CO_2]_{vb} = \left( M_{M_b} / M_g \right) \times 10^6,
\]

where \( M_{M_b} \) is the mass of glass within the MI, calculated as the glass volume multiplied by a melt density that is assumed to be 2.75 g cm⁻³ [46]. The results show that bubbles in MIs from the lamprophyres contain 323 to 47,940 ppm CO₂ (N = 29), and >93% of bubbles have a CO₂ content of >1000 ppm (0.1 wt%) (Fig. 5f, Table S6). The calculated CO₂ concentrations in MIs of the lamprophyres range from 474 to 47,641 ppm (N = 29), with most (>80%) higher than 5000 ppm. Silicate crystal-hosted MIs, representing melts during various stages of an evolving magmatic system, can be analyzed to constrain the CO₂ contents that dissolved in the melt before volcanic eruption and/or degassing [47]. We thus estimate that the measured CO₂ concentrations represent those of the pre-eruptive and possibly evolved lamprophyre magmas, which mostly fall between 0.5 wt% and 2.0 wt%. These contents are similar to or even higher than the CO₂ concentrations (0.5–1.0 wt%) in MIs of the end-Triassic Central Atlantic Magmatic Province basalts, which were estimated by the same method [48]. It should be noted that a high CO₂ concentration of MIs is mainly observed in High-MgO lamprophyres with light δ²⁶Mg values (see Supplementary Data), and the number of MIs in High-MgO lamprophyres is much larger than that in Low-MgO lamprophyres. This probably indicates that High-MgO lamprophyres with recycled carbonates in their mantle sources contain more MIs and higher CO₂ concentrations in the pre-eruptive magmas, although low-volume melts could also have extremely high CO₂ content even if the source is not specifically C rich, due to the strong incompatibility of CO₂ in peridotite [49]. Because most of the early Cretaceous lamprophyres in the eastern NCC belong to the High-MgO group with low δ²⁶Mg (Fig. 2), their sources were plausibly most strongly affected by carbonate metasomatism and attendant enrichment in carbon. Melting of this metasomatized CLM then produced primary magmas with high CO₂ content, which may have been further enhanced
CO2 outgassing is thus expected during the period represented by lamprophyres. A larger flux of the eastern NCC are much more abundant than those represented by lamprophyres. Thus, early Cretaceous mantle-derived magmas in the eastern NCC are much more abundant than those represented by lamprophyres. A larger flux of CO2 outgassing is thus expected during the period of extensive destruction of the NCC, in addition to lamprophyres. Intrusion or eruption of these magmas could have carried a large amount of CO2 from the mantle into the surface.

Experimental studies show that the solubility of carbon dioxide in melts decreases at lower pressures, and CO2 can even be directly degassed at mantle depths [50]. Therefore, the thinned lithosphere, as a result of cratonic destruction and extension, can further facilitate CO2 outgassing via magmatism. There is no evidence for the presence of a deep-sourced mantle plume beneath the eastern NCC during the Phanerozoic era. We thus propose that the destruction of the CLM represents another important cause of CO2 emission from the mantle, in addition to continental rifting, active island arc volcanism and mantle plume. During this process, carbon in the CLM can experience gradual oxidation during mantle upwelling, with a change of carbon speciation from a reduced to an oxidized form, and a portion of carbon can be liberated via redox melting from the reduced mantle [11].

Enormous CO2 reservoirs can be formed by the eruption or intrusion of magmas. There are abundant crustal CO2 reservoirs in the Songliao and Bohai Bay basins in the eastern NCC, which indicates that the volume of CO2 degassing is enormous [51]. CO2 reservoirs in the Songliao basin were formed primarily in Cretaceous, and voluminous inorganic CO2 (mainly mantle-derived and crust-derived) is observed in these reservoirs. For example, the high CO2 content (>90%) and δ13C (−4.95‰) and high helium isotopic composition (R/Ra = 3.34) of Wanjinta reservoirs indicate that the CO2 was chiefly sourced from the mantle [51]. The Bohai Bay basin, a Mesozoic-Cenozoic basin, is the central area of destruction of the eastern NCC. The reservoirs there also have high CO2 content (79.17–98.61%) and high R/Ra (2–3.34), which indicates that the CO2 was derived from the mantle [51].

A recent study by Aulbach et al. [8] quantified the CO2 flux related to the reaction of the CLM with silica-undersaturated (carbonated) melt, referred to as wehrlitization, and linked this flux to surficial degassing in rifts and basins. As discussed above, abundant wehrlite and pyroxenite xenoliths are found in Mesozoic mantle-derived rocks in the eastern NCC, and many of these xenoliths have light Mg isotopic compositions and high Ca/Al ratios (Fig. 3). For instance, the characteristics of low Ti/Eu, high Ca/Al, (La/Yb)N and Zr/Hf of clinopyroxenes in Liaoyuan wehrlites are ascribed to interaction with a silicic ore. Decarbonation of these melts is initially out of thermal and compositional equilibrium, causing intensive melt-rock reactions. During this process, the following reaction will happen at ~1.5–2.0 GPa [52]:

\[
\text{Enstatite + dolomite (melt) = forsterite + diopside + CO2 (vapor)}.
\]

A quantitative estimate suggests that 2.9 to 10.2 kg CO2 can be released per 100 kg of wehrlite formed [8]. Extensive CO2 release is suggested to have occurred during the carbonated melt–CLM reaction process in the course of this destruction of the eastern NCC, along with the Tan-Lu Fault Belt, which was most active in the early Cretaceous [33].

Stable continents are long-term storage sites for sedimentary carbonates, and the amount of carbonates stored in continents is thought to be at least 10 times greater than that stored in oceanic crust [53]. Carbonates in crusts can be trapped by plutons that ascend to shallow levels in the arc crust or are transported into the lower crust during later arc stages. Global flare-ups in continental arc volcanism were proposed to have the potential to release CO2 as a result of magmatic interaction with ancient crustal carbonates stored in the continental crust [54,55]. The eastern NCC is typically characterized by a giant felsic magmatism event at the early Cretaceous with a volume much larger than that of mafic magmatism [17], implying large-scale crustal melting and reworking during the cratonic destruction process. These early Cretaceous felsic magmas (i.e. granites) have high zirconium saturation temperatures and contain an important contribution from the hot upwelling mantle [25]. Decarbonation is expected to widely occur during interaction between the hot felsic magmas and the limestones chronically stored in the continental crust. This process could also contribute to CO2 release, in addition to mantle CO2 outgassing via mafic magmatism and carbonated melt–CLM reaction during destruction of the eastern NCC (Fig. 6).
POSSIBLE CONTRIBUTIONS TO THE CRETACEOUS GREENHOUSE

The amount of CO₂ outgassing induced by the destruction of the eastern NCC could be significant, particularly if one considers that the removed CLM contained a large amount of recycled carbon from subducted slabs prior to thinning. The fluxes of deep CO₂ outgassing during this destruction process may be large given the short duration of mantle-derived, CO₂-rich magmatism (both intrusions and volcanics; Fig. 1) in the early Cretaceous (∼125 Ma) as well as the strong carbonated silicate melt–CLM reaction that resulted in substantial CO₂ release along the massive Tan-Lu Fault Belt [8]. At a larger scale, enormous quantities of CO₂ that were rapidly released into the atmosphere, induced by the destruction, may have perturbed the global climate and partly contributed to the atmospheric CO₂ rise during the Cretaceous, one of the longest greenhouse periods of Earth’s history, with atmospheric CO₂ levels 4 to 10 times higher than those prior to the Industrial Revolution [56].

CONCLUSIONS

We present the first Mg isotope data for early Cretaceous lamprophyres and collate available chemical and Mg isotopic data for mantle xenoliths in Paleozoic diamond-bearing kimberlites and Mesozoic mafic igneous rocks as well as orogenic ultramafic massifs in the NCC. These results suggest the presence of a widespread, C-rich CLM beneath the NCC at and before lithospheric thinning in the early Cretaceous, with an important contribution from recycled carbonate sediments. Long-term and threesided—i.e. the south, north and east—oceanic plate subductions underneath the NCC during the Paleozoic and Mesozoic could have contributed a vast amount of carbon to the lithospheric mantle of the NCC. Redox melting of the reduced, C-rich CLM as it was exhumed to shallower depths due to lithospheric extension and thinning in the early Cretaceous generated large amounts of basaltic lavas and lamprophyres, resulting in the release of voluminous CO₂ to the exosphere. This may represent an important cause of CO₂ emission from the mantle, in addition to mantle plumes, active island arc volcanism and continental rifts, as proposed in previous studies [1,3–7]. The amount of magmatic CO₂ outgassing is largely supplemented by the release of mantle CO₂ induced by the carbonated melt–CLM reaction [8] and decarbonation induced by interaction between hot felsic magmas and crustal limestones. Therefore, deep CO₂ outgassing can be linked to the destruction of a long-term stable craton and can be said to have enhanced global CO₂ input into the atmosphere.

SUPPLEMENTARY DATA

Supplementary data are available at NSR online.

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AUTHOR CONTRIBUTIONS

S.-A.L. and S.G.L. designed the project. Z.-X.W. and D.L. analyzed all data. S.-A.L. and Z.X.W. developed the manuscript with contributions from other co-authors.

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