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# Estimating the carbon content of the deep mantle with Icelandic melt inclusions



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#### A R T I C L E I N F O

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# ABSTRACT

Earth's carbon budget is central to our understanding of the long-term co-evolution of life and the planet. Direct observations of surface reservoirs allow for the detailed quantification of their carbon content. However, the carbon content of Earth's deep interior remains poorly constrained. Here we study olivinehosted melt inclusions from two Icelandic eruptions, with those from the Miðfell eruption allowing us to investigate the carbon content of the deep mantle. Comparison with the previously studied Borgarhraun eruption highlights the presence of deep, plume-sourced mantle material within the Midfell source region. Miðfell contains trace element-depleted melt inclusions undersaturated in CO<sub>2</sub>, which have high  $CO_2/Ba$  (= 396 ± 48) and  $CO_2/Nb$  (= 1832 ± 316), though some inclusions preserve even greater relative carbon enrichment. These observations allow us to reconstruct the CO<sub>2</sub> content of the bulk Miðfell source as being > 690 ppm. By identifying that Miðfell is a mixture of depleted and deep mantle components, we can estimate a CO<sub>2</sub> content for the deep mantle component of  $1350 \pm 350$  ppm; a concentration that is over ten times higher than depleted MORB mantle estimates. Assuming that the deep mantle component identified in Midfell is representative of a global reservoir, then with our new CO<sub>2</sub> estimate and by considering a range of representative mantle fractions for this reservoir, we calculate that it contains up to 14 times more carbon than that of the atmosphere, oceans, and crust combined. Our result of elevated CO<sub>2</sub>/Ba and CO<sub>2</sub>/Nb ratios, and carbon enrichment support geochemical bulk Earth carbon models that call for the presence of carbon-rich deep mantle domains to balance Earth's relatively carbon-poor upper mantle and surface environment.

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# 1. Introduction

Geological processes have modulated the Earth's atmospheric carbon content for billions of years (Hayes and Waldbauer, 2006). At the heart of this cycle is a partitioning of carbon between planetary reservoirs, one that has placed the overwhelming majority of carbon in Earth's crust, mantle, and core (e.g., Dasgupta and Hirschmann, 2010). The carbon flux from these solid-Earth reservoirs to the atmosphere is efficiently returned via silicate weathering (e.g., Walker et al., 1981), maintaining a habitable climate, and closing the loop on a cycle that has helped maintain liquid water at Earth's surface over almost its entire history (Mojzsis et al., 2001).

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To understand how Earth has come to operate such a stable and long-lasting chemical cycle, it is key to know how carbon is distributed among its reservoirs (Hirschmann, 2016). Whilst for the atmosphere, oceans, and to some extent the crust, their carbon content can be measured directly, quantifying the carbon content of mantle reservoirs is more challenging. Two basic problems frustrate accurate estimation of mantle carbon content: (i) the low solubility of carbon in basaltic melts (Shishkina et al., 2010), which means that information on high-carbon mantle domains is preferentially lost as their melts begin to degas at high pressure in the crust or shallow mantle; and (ii) the partial view that volcanism provides of the mantle, with the vast majority of magmas tapping the depleted upper mantle. Only rarely do volcanics display evidence for the involvement of deep-sourced material in their petrogenesis: such occurrences are usually linked to the presence of mantle plumes. These compounding problems cause considerable uncertainty in previously published mantle carbon estimates.

In this paper we present new observations that help constrain the amount of carbon in the deep mantle reservoir. In Section 2, we review the geochemical constraints available on mantle carbon, and identify the observational gaps in our present reservoir inventories. We next present our methods (Section 3) and new data (Section 4) from two Icelandic eruptions, demonstrating that their geochemical characteristics are suited to quantify the amount of carbon in the deep mantle. We show that one of these eruptions, Miðfell, contains melt inclusions that are undegassed and relatively enriched in carbon compared to lithophile trace elements of similar mineral-melt compatibility (Section 5). In Section 6, we use these results to place a new constraint on the deep mantle carbon content, and in Section 7 discuss the implications of this constraint for the origin and distribution of carbon in the Earth.

# 2. Measuring mantle carbon

An enormous amount of work, especially over the last two decades, has focused on measuring the carbon content of mantlederived volcanics. Despite this effort, surprisingly few observations provide tight constraints on upper mantle carbon content, and fewer still on deep mantle carbon. This section aims to contextualise the present study with this body of work, and identify the features that, in subsequent sections, will mark our new observations as distinct from pre-existing data sets.

# 2.1. The upper mantle

Mid-ocean ridge basalts (MORBs) sample the most accessible mantle reservoir, the depleted MORB mantle (DMM). Although a shallow reservoir in the context of the mantle, the DMM can be a component of deep-sourced mantle plume magmas, as is the case on Iceland (Stracke, 2012).

A key technique for estimating carbon in DMM is to find an incompatible lithophile trace element (ITE) to reference MORB carbon concentrations against, thereby providing a control for fractionation during mantle melting and crustal differentiation. Estimates of source mantle ITE concentration (e.g., Workman and Hart, 2005) then enable calculation of a source carbon content. Ba and Nb are commonly chosen as reference elements because they have similar compatibility to carbon during peridotite melting (Rosenthal et al., 2015). However, as Ba and Nb are not volatile, accurate estimation of source carbon is dependent on either finding melts that are CO<sub>2</sub> undersaturated, or reconstructing pre-degassed CO<sub>2</sub>. A degassing correction has been applied to some highly vesicular glass samples, which re-coupled CO<sub>2</sub> to ITE concentrations, giving DMM carbon estimates of  $393 \pm 82$  ppm CO<sub>2</sub> using CO<sub>2</sub>/Nb = 534, and  $427 \pm 45$  ppm CO<sub>2</sub> using CO<sub>2</sub>/Ba = 106 (Table 1; 14°N Mid-Atlantic Ridge; Cartigny et al., 2008). We note that whilst carbon exists in the mantle in oxidised and reduced forms (Dasgupta and Hirschmann, 2010), for ease of comparison between data sets, we report total carbon as CO<sub>2</sub>.

Only rare suites, such as some pillow glasses erupted at relatively high pressure and with intrinsically low carbon content, may show undersaturated volatile concentrations. Michael and Graham (2015) used such samples to estimate a DMM CO<sub>2</sub> of  $59 \pm 39$  ppm (Table 1; CO<sub>2</sub>/Ba = 105; Global MORB). A difficulty with this approach is in having confidence that measured melts are truly undegassed. Saal et al. (2002) emphasised how linearly correlated carbon-trace element concentrations can validate a data set as having undersaturated melt populations. However, Matthews et al. (2017) presented a forward model to suggest that such correlations can readily arise from the mixing of variably degassed melts at low pressure. They concluded that to have more confidence in identifying undegassed melts, data will ideally show some melts with high

Table 1								
Estimates of mantle carbc	on reservoirs.	This study pre	esents carbon estimates for l	Miðfell source mantle and a deep ma	antle reservoir, calculated fro	m olivine-hosted melt in	clusion compositions and a I	DM-Deep mixing model. CO <sub>2</sub> /ITE
ratios for each data set al element concentrations.	re as publishe	d in their orig	ginal publications. Source tra	ice element and $CO_2$ estimates are the theorem of the second s	ne values presented by Hauri	i et al. (2018), who used	the method of Shimizu et al.	(2016) to calculate source trace
Setting	Ocean Isla	nd			Mid Ocean Ridge			
Locality Reference	Miðfell This study		Borgarhraun Hauri et al. (2018)	Hawaii Anderson and Poland (2017)	Eq. MAR Le Voyer et al. (2017)	Siquieros Saal et al. (2002)	14°N MAR Cartigny et al. (2008)	Global MORB Michael and Graham (2015)
	Bulk	Deep						
CO <sub>2</sub> /Nb	1832	> 1520	391	758	557	230	534	607
CO <sub>2</sub> /Ba	396	> 150	48.3	86	96.6	74.2	106	105
Mantle CO <sub>2</sub> (Nb)	887	I	129	1	48	23	393	06
Mantle CO <sub>2</sub> (Ba)	1580	I	80	1	22	24	427	59
Mantle CO <sub>2</sub> (other)	> 690	> 1000	1	263	I	I	1	1
Nb ppm	0.484	0.658	0.330	0.347	0.0862	0.100	0.736	0.149
Ba ppm	3.99	6.60	1.66	3.06	0.228	0.323	4.04	0.562

ITE content that have clearly degassed to their saturation concentration (i.e., no ITE-carbon correlation), and others with lower ITE and carbon content where an ITE-carbon correlation persists; such suites evidence incomplete melt mixing. Data sets exhibiting such structure offer the best chance for empirically separating degassed and undegassed melt populations.

Applying the insights from Matthews et al. (2017) can make certain data sets difficult to empirically validate for carbon undersaturation. For example, observations of quenched matrix glasses, for which the melt has been largely homogenised prior to eruption (i.e., many seafloor basalts; Shorttle, 2015; Michael and Graham, 2015), do not provide data sets that can be internally validated: giving only one CO<sub>2</sub> and ITE content per eruption. One solution is to use olivine-hosted melt inclusions, which, when trapped at high pressure, may capture melts with both undegassed carbon and chemical diversity that has not been homogenised by mixing (Maclennan, 2008b). Melt inclusion studies have thus been key in characterising upper mantle carbon: their results suggest that the DMM contains carbon heterogeneity, as both CO<sub>2</sub>/ITE ratios and estimated ITE concentrations vary (Le Voyer et al., 2017), with carbon concentrations ranging from 22-427 ppm CO<sub>2</sub> (Table 1; Sigueiros, Saal et al., 2002; Mid-Atlantic Ridge, Cartigny et al., 2008, Le Voyer et al., 2017).

# 2.2. The deep mantle

We use the term 'deep mantle' to refer loosely to the region of Earth's convecting mantle that is not represented by the composition of typical MORB. The geometry of this reservoir is poorly constrained, and its upper horizon could fall anywhere from the mantle transition zone, to the large low shear velocity provinces above the core-mantle boundary (e.g., Hofmann, 1997). Upwelling plumes can bring this deep mantle material into the upper mantle and melt it at shallow levels. Whatever its locus, the deep mantle appears to comprise primitive material that has remained isolated for billions of years (e.g., Mukhopadhyay, 2012; Peters et al., 2018), as well as crustal material introduced via recycling processes (e.g., Nestola et al., 2018). Both components could be significant carbon reservoirs: primitive material potentially containing solar nebular carbon, which dissolved into Earth's early magma ocean along with noble gases (e.g., Williams and Mukhopadhyay, 2018); whereas, recycled material could introduce organic or inorganic carbon from the surface (e.g., Nestola et al., 2018).

Ocean island basalts (OIBs) are prime candidates in the search for deep mantle carbon. Geochemical observations have consistently shown OIB to have more evidence of both recycled and primitive mantle components than MORB (e.g., Hofmann, 1997), and in many cases geophysical observations support their lower mantle origins (e.g., Montelli et al., 2006). Beneath Iceland specifically, the hot, low velocity, plume conduit has been seismically imaged through the deflected transition zone (Jenkins et al., 2016), and down into the lower mantle (Yuan and Romanowicz, 2017).

The nature of magma generation and eruption at ocean island settings typically results in OIB degassing (Gonnermann and Mukhopadhyay, 2007): low eruption or melt inclusion entrapment pressures combine with high initial dissolved CO<sub>2</sub> content, derived from low degree melting and/or carbon-rich source material. This degassing can, however, be an advantage if rather than measuring the melt, the degassed CO<sub>2</sub> flux is measured. Anderson and Poland (2017) measured CO<sub>2</sub> degassing at Hawaii, along with a melt production rate to estimate the Hawaiian source mantle carbon content to be 962 + 296/ - 227 ppm CO<sub>2</sub>. This estimate is significantly higher than that obtained from MORB (Table 1), a discrepancy suggesting the presence of high carbon regions in Earth's deep mantle. However, bulk degassing cannot be used to uniquely assign carbon to the recycled (e.g., Sobolev et al., 2005) or primitive (e.g., Trieloff et al., 2000) components inferred for the Hawaiian source. Our approach minimises this ambiguity by focusing on melt inclusions from single eruptions.

#### 2.3. Bulk silicate Earth carbon estimates

In principle, the size of the bulk silicate Earth (BSE) carbon reservoir could be reconstructed using carbon estimates of all mantle reservoirs, including the deep mantle. However, as emphasised above, a key piece of information is poorly known: the size of this deep mantle reservoir (Hofmann, 1997). A common approach to this problem is to assume, or infer from geochemical mass balance, the size of the deep mantle reservoir and then use carbon-gas or carbon-trace element ratios to extrapolate to bulk Earth.

A recent review paper from Halliday (2013) presented several models for estimating bulk Earth carbon content. The 'basalt' model used the water content of MORB and OIB to infer a bulk water content for the entire mantle, and then took representative H/C ratios to calculate a bulk mantle carbon content (163 ppm CO<sub>2</sub>). The 'layered mantle' model used Ar isotope budgets across all Earth's reservoirs, along with volatile ratios, to derive a bulk Earth carbon budget of 2462 ppm CO<sub>2</sub>. This estimate is very similar to the value from Marty (2012), who also used an <sup>40</sup>Ar budget to extrapolate observed C/N and C/<sup>4</sup>He ratios to the whole Earth (2831 ppm CO<sub>2</sub>).

Carbon-ITE ratios in MORB, enriched-MORB, and OIB have also been used to reconstruct BSE carbon. Hirschmann (2018) presented a recent implementation of this approach, taking  $CO_2/Ba =$  $100\pm20$  for the convecting mantle and a representative Ba concentration to estimate  $514\pm147$  ppm  $CO_2$  in BSE. This new estimate assumes that oceanic basalts, regardless of enrichment, preserve similar  $CO_2/Ba$  ratios, and that analysed oceanic basalts sample all significant geochemical reservoirs within the Earth. Our new results will show, in contrast, that there is evidence for both significant  $CO_2/Ba$  variability, and that there exist mantle reservoirs with far higher  $CO_2/Ba$  than have previously been identified in MORB or OIB.

Whilst the above BSE carbon estimates vary significantly, they have a common implication: assuming DMM carbon concentrations of 20–100 ppm CO<sub>2</sub> (e.g., Dasgupta and Hirschmann, 2010), a significantly more carbon-rich reservoir must exist in the deep Earth to balance high BSE carbon estimates. For example, Hirschmann (2016) calculated that this reservoir needs  $CO_2/Nb = 5000 \pm 2000$  and  $CO_2/Ba = 850 \pm 350$  to match the BSE carbon content calculated by Marty (2012).

#### 2.4. Carbon in the Icelandic mantle

Previous studies of Icelandic geothermal fluids, glasses and vesicles have concluded that the Icelandic mantle source has little to no carbon enrichment with respect to DMM. While these studies used bulk analyses to assess average source carbon, our study uses olivine-hosted melt inclusions to access signals of source heterogeneity that are preserved in melts prior to mixing.

Only one melt inclusion suite from Iceland has previously been interrogated for mantle carbon signatures; the Borgarhraun eruption in the Northern Rift Zone (NRZ, Fig. 1a). Olivine-hosted melt inclusions from Borgarhraun record compositional heterogeneity (Maclennan et al., 2003), show evidence of concurrent crystallisation and melt mixing (Maclennan, 2008a), and preserve a CO<sub>2</sub>-ITE correlation (Hauri et al., 2018), which has been used as evidence for an undegassed CO<sub>2</sub> signature (CO<sub>2</sub>/Ba =  $48.3 \pm 2.7$ , CO<sub>2</sub>/Nb =  $391 \pm 16$ ; Table 1). The mantle carbon content inferred from these observations is similar to that of DMM, with CO<sub>2</sub>/Nb comparable



**Fig. 1.** Icelandic eruptions discussed in this study: Miðfell (red), Kistufell (purple), and Borgarhraun (blue). Map of Iceland in (a) shows eruption locations, central volcanoes (red areas), and main rift zones (orange): WRZ, western rift zone; ERZ, eastern rift zone; and NRZ, northern rift zone. In (b) He isotope compositions relative to atmospheric  ${}^{3}$ He/<sup>4</sup>He (R/R<sub>a</sub>) and in (c) Sr isotope compositions against  $\epsilon$ Nd for key eruptions (diamonds) and young Icelandic eruptions (< 120 kyrs) from rift zones (grey). Kistufell and Miðfell show elevated  ${}^{3}$ He/<sup>4</sup>He with respect to Borgarhraun. Data compiled in Shorttle and Maclennan (2011). (For interpretation of the colours in the figures, the reader is referred to the web version of this article.)

to undegassed MORB suites (Table 1), but with  $CO_2/Ba$  half the inferred global average (Hirschmann, 2018).

Borgarhraun melts are not strongly enriched in lithophile elements, being depleted relative to other Icelandic melts, and its He isotope signature  $(R/R_a = 7.9; Fig. 1b)$  lies within the MORB range (Füri et al., 2010). Hauri et al. (2018) suggested that the Borgarhraun mantle contains a source common to the Atlantic depleted mantle: hydrated and carbonated material originating from subduction-related modification. These features strongly suggest that there is no deep mantle component in the Borgarhraun source, which fits the regional systematics of 'MORB-like' He-Ne in NRZ eruptions in contrast to more 'solar-like' noble gas compositions of Western Rift Zone (WRZ) eruptions (e.g., Füri et al., 2010). Such observations align with Pb isotope constraints indicating distinct mantle source components across Iceland (Shorttle et al., 2013), and merit investigation of whether there are associated differences in mantle carbon abundances between the two rift zones. Our new observations from eruptions in the WRZ and central Iceland (Fig. 1a) directly address this question.

# 3. Samples and methods

# 3.1. Geological context

We present new data from two Icelandic eruptions, Miðfell (also known as Dagmálafell;  $64^{\circ}$  10.456' N,  $021^{\circ}$  02.859' W) and Kistufell ( $64^{\circ}$  47.442' N,  $017^{\circ}$  10.456' W; Fig. 1a). Miðfell is a  $\sim$ 300 m high mountain in Iceland's WRZ, lying on the eastern edge of Þingvallavatn and within the Hengill volcanic system. It is a northeast-southwest striking ridge, which is thought to have erupted beneath ice during the last glacial period (Gurenko and Chaussidon, 1995, and references therein). The mountain's lower flanks comprise vesicular, olivine-phyric, pillow basalts with glassy rinds, whereas the upper portions are composed of hyaloclastite. Kistufell is a table mountain located in central Iceland, at the northern edge of Vatnajökull. Kistufell may have erupted towards the end of the last glacial period as the ice-sheet retreated (Bred-

dam, 2002). Glassy pillows are exposed at the northern flanks of the mountain.

# 3.2. Geochemical context

Previous Miðfell melt inclusion studies have measured significant trace element variability, including high-degree melts with diluted ITE concentrations (Gurenko and Chaussidon, 1995). If carbon remained coupled to these ITE's, then it is likely that some Miðfell melt inclusions have avoided degassing, providing an opportunity to recover mantle carbon content. Both Miðfell and Kistufell have noble gas isotope data that support the presence of primordial material in their source regions:  $R/R_a \sim 17$  (Breddam et al., 2000; Füri et al., 2010) and, where heavy noble gases have been measured in Miðfell, primordial Xe and Ne isotopic ratios (Mukhopadhyay, 2012). Miðfell olivines and matrix glass also show noble gas evidence for other components, including recycled air (e.g., Harrison et al., 1999). Combined, these observations make Miðfell and Kistufell good targets for identifying deep mantle carbon.

#### 3.3. Analytical methods

Olivine phenocrysts 0.5-4 mm in size were picked from hand-crushed pillow glasses. Care was taken to pick unaltered olivine macrocrysts containing glassy melt inclusions without postentrapment crystals. Olivines were individually mounted and polished to expose melt inclusions before being re-mounted and polished for analysis. Melt inclusions were analysed for trace elements, CO<sub>2</sub>, and H<sub>2</sub>O by secondary ion mass spectroscopy at the Edinburgh ion microprobe facility. Major elements were measured by electron probe microanalysis. A subset of the Miðfell melt inclusion suite was processed at Woods Hole Oceanographic Institution to reconstruct their initial CO<sub>2</sub> content from vapour bubble and inclusion glass CO<sub>2</sub> concentrations. Melt inclusion and vapour bubble volumes were determined by X-ray tomography. Vapour bubble CO<sub>2</sub> content was determined using confocal Raman spectroscopy.



**Fig. 2.** (a) La/Yb ratio of Miðfell (red) and Kistufell (purple) melt inclusions against host olivine forsterite content. Average melt inclusion La/Yb ratios are shown by horizontal lines, while matrix glass averages are indicated by arrowheads at  $X_{Fo}^{OI} = 0.95$ . The olivine forsterite compositions in equilibrium with the matrix glasses are shown by inverted triangles at La/Yb = 8. Melt inclusion variability is preserved with decreasing forsterite content in the Miðfell suite, while the Kistufell melt inclusions have much lower La/Yb variability and are clustered within a smaller forsterite content range. (b) CO<sub>2</sub>/Ba against forsterite, indicating that the highest CO<sub>2</sub>/Ba ratios are preserved in the most forsteritic olivines. One sigma error ellipse plotted in grey.

For full method details and representative melt inclusion micrographs, see Supplementary Material.

#### 4. Results

### 4.1. Major elements

Miðfell olivine phenocryst compositions range from  $Fo_{90.9}$  to  $Fo_{85.2}$  (Fig. 2), the matrix glass and olivine-hosted melt inclusion compositions are basaltic (SiO<sub>2</sub> ~48 wt%), giving a picritic whole rock composition (Gurenko and Chaussidon, 1995). Kistufell olivines range from  $Fo_{89.7}$  to  $Fo_{87.9}$  with basaltic melt inclusions (SiO<sub>2</sub> ~48 wt%).

Melt inclusion major element compositions have been corrected for post-entrapment crystallisation by iteratively adding an equilibrium olivine composition, assuming an Fe-Mg K<sub>D</sub> = 0.34 (Matzen et al., 2011), until the melt inclusion reached equilibrium with its host (Danyushevsky et al., 2000). For Miðfell, this correction takes into account the ferric iron content of each melt inclusion (see Supplementary Material), whereas for Kistufell we assumed a constant ferric to total iron ratio of 0.07 based on matrix glass measurements (Breddam, 2002). The majority of melt inclusions required < 5% olivine addition to bring them back into Fe-Mg equilibrium with their olivine host.

# 4.2. Trace elements

Miðfell melt inclusions record substantially more variability than those from Kistufell (Fig. 2 & 3). The method of Maclennan et al. (2003) was used to establish that the Miðfell analyses for all major, trace, and volatile elements (apart from H<sub>2</sub>O, Yb and Lu) have a signal-to-noise ratio > 1, and > 99% confidence that natural variability can be resolved from analytical noise. These inclusions preserve trace element variability comparable with that found by the combination of all previously published undegassed melt inclusion and MORB glass suites (e.g., Saal et al., 2002). Kistufell melt inclusions recover a signal-to-noise ratio > 1 for the majority of major, trace, and volatile elements. However, the Kistufell melt inclusion suite has much lower melt heterogeneity than Miðfell, as indicated by their contrasting signal-to-noise ratios (e.g., Miðfell  $\sigma_t/\sigma_r|_{Ba} = 3.06$ ).

Multi-element plots (Fig. 3) illustrate Miðfell and Kistufell trace element variability compared to that of the Borgarhraun eruption (Hauri et al., 2018). As is typical for many mantle-derived basalts, the most incompatible trace elements show the largest concentration range. In Miðfell, Ba content ranges from 0.37–115 ppm

(29.8–3.5% relative error; matrix glass 6.3 ppm), and Nb content ranges from 0.04–22.9 ppm (17.6–9.1% relative error; matrix glass 0.78 ppm). Vapour bubbles only occur in Miðfell melt inclusions with high trace element concentrations (> 12.4 ppm Ba and > 2.47 ppm Nb). Kistufell melt inclusions rarely contain vapour bubbles, so no inclusions with vapour bubbles were measured for this data set. Trace element patterns in all three eruptions show that some of the melt inclusions preserve positive Sr and negative Zr anomalies, consistent with previous Miðfell melt inclusion analyses (Gurenko and Chaussidon, 1995).

#### 4.3. Volatile elements

Kistufell melt inclusions have a water content that matches that of the matrix glass surrounding host olivines, but the sample population shows some variability (0.10–0.19 wt%). Water content in the Miðfell melt inclusions and matrix glass is almost constant at ~0.06 wt%, which is low compared to other Icelandic melt inclusion suites (e.g., Laki, ~0.65 wt% H<sub>2</sub>O, Hartley et al., 2015). Melt inclusion CO<sub>2</sub> content varies from 20–1120 ppm (Fig. 4a). At low trace element concentrations (Ba < 3.0 ppm, Nb < 0.2 ppm) the maximum melt inclusion CO<sub>2</sub> content is controlled by CO<sub>2</sub>/ITE ratios. At higher trace element concentrations, maximum CO<sub>2</sub> content reaches a plateau at 1120 ppm. Melt inclusions containing vapour bubbles record a glass CO<sub>2</sub> content of 720–1170 ppm, consistent with the vapour bubble-absent inclusions, and a total (reconstructed) CO<sub>2</sub> content of 1340–4550 ppm (Fig. 4a). Matrix glass has a CO<sub>2</sub> content of < 200 ppm.

#### 4.4. CO<sub>2</sub>/ITE ratios

Kistufell melt inclusions record maximum  $CO_2/Nb = 391 \pm 70$ and  $CO_2/Ba = 71.9 \pm 13.9$ , which are associated with the lowest ITE concentration melts. Miðfell melt inclusions preserve maximum  $CO_2/Nb = 5737 \pm 987$  and  $CO_2/Ba = 566 \pm 68$ , also associated with low trace element concentrations, i.e., lower than the matrix glass (Fig. 5), and hosted in the most forsterite-rich olivines (Fig. 2b). Melt inclusions with  $CO_2$  content reconstructed from vapour bubble concentrations also have high  $CO_2/ITE$  ratios, with maximum  $CO_2/Nb = 1186 \pm 127$  and  $CO_2/Ba = 236 \pm 25$ . Comparison to previously analysed melt inclusion and MORB glass suites shows that the Miðfell melt inclusions preserve some of the highest  $CO_2/ITE$  ratios ever measured in natural basaltic glass (e.g., Le Voyer et al., 2017). Importantly, Miðfell inclusions exhibit these higher  $CO_2/ITE$  ratios at the same ITE concentrations as in



**Fig. 3.** Trace element spider diagrams showing compositional variability in (a) Miðfell, (b) Kistufell, and (c) Borgarhraun (Hauri et al., 2018) melt inclusion suites, normalised to primitive mantle (PM; BSE composition, McDonough and Sun, 1995). Inclusions with  $CO_2/Ba > 150$  are coloured, the rest are grey (Borgarhraun melt inclusions have an average  $CO_2/Ba \sim 48$ ). Matrix glass compositions are shown as dark squares. The likely position of carbon, given its measured compatibility (Rosenthal et al., 2015), is shown as a blue vertical line. In (c) the light red line and circles are the Miðfell average melt inclusion composition, and in light blue is the Borgarhraun melt inclusion average. Miðfell and Borgarhraun matrix glasses are very similar, but Miðfell shows greater trace element variability within the melt inclusion population.

previously studied suites, indicating that their relative carbon enrichment is not a feature of anomalous trace element depletion or enrichment (Fig. 5).

#### 5. Does Miðfell preserve a mantle carbon signature?

The previous section demonstrated that whilst Miðfell melt inclusions preserve significant trace element variability, Kistufell contains a more homogeneous melt inclusion population. On this basis, the Kistufell melt inclusions do not enable empirical validation of whether they have degassed carbon (Fig. 4b; Matthews et al., 2017). However, Kistufell melt inclusions have ITE concentrations that in the Miðfell melt inclusion population are associated with carbon loss (Fig. 4). For Miðfell, this carbon loss is shown by systematically decreasing  $CO_2/ITE$  ratios with increasing ITE concentration (Fig. 5c,d). Kistufell inclusions have likely degassed, therefore we focus our investigation of deep mantle carbon onto the Miðfell inclusions, where we can be sure of recovering undegassed systematics.

Trace element and  $CO_2$  data from Miðfell melt inclusions suggest that some process has enriched these basaltic glasses in carbon, relative to ITE's, compared with other inclusion and glass suites (e.g., Michael and Graham, 2015). To use this observation to constrain the carbon content of the Miðfell mantle source first requires an assessment of the influence of crustal processes, which could have affected the carbon and ITE content of Miðfell melts.

# 5.1. Degassing and olivine decrepitation

The strongest signal in the Miðfell CO<sub>2</sub> data, seen most clearly at trace element concentrations greater than that of the matrix glass, is one of degassing (Fig. 4a). At these high ITE concentrations, melt inclusions with variable trace element concentrations have constant CO<sub>2</sub> content, indicating the partial loss of initial CO<sub>2</sub>, and therefore loss of information on carbon from deeper in the system. If degassing occurred in some of the Miðfell melts prior to entrapment, then solubility models suggest entrapment pressures of ~1.8 kbar (taking 1170 ppm CO<sub>2</sub>; Shishkina et al., 2010).

However, the presence of vapour bubbles in some of the trace element enriched melt inclusions suggests that some Miðfell melts were initially trapped with higher dissolved CO<sub>2</sub> concentrations and underwent exsolution post-entrapment. Reconstructed CO<sub>2</sub> concentrations from these inclusions are consistent with saturation pressures of up to ~4.5 kbar (~13 km depth). If all melts started with the same CO<sub>2</sub>/ITE ratio, then those CO<sub>2</sub>-saturated melt inclusions lacking vapour bubbles must have lost exsolved CO<sub>2</sub> during ascent to eruption, perhaps by decrepitation (Maclennan, 2017).

ITE-rich Miðfell melts appear to have been degassed to their saturation concentration, but at lower ITE concentrations melts show an ITE-CO<sub>2</sub> correlation. This observation suggests that ITE-depleted melt inclusions preserve an undegassed CO<sub>2</sub> signal (Matthews et al., 2017); an inference supported by bubble-reconstructed melt inclusions that have CO<sub>2</sub>/ITE ratios in broad agreement with those of depleted inclusions (Fig. 4 & 5). Therefore, relative carbon enrichment in Miðfell was likely not restricted to the most ITE-depleted melts, but rather a characteristic of all ITE concentrations (Fig. 4a & 5). With this result, the key question is what source or process led to high CO<sub>2</sub>/ITE ratios in Miðfell inclusions?

#### 5.2. Crustal melt modification

Gurenko and Sobolev (2006) analysed olivine-hosted melt inclusions and gabbroic xenoliths from Miðfell to infer that the trace element chemistry of Miðfell melt inclusions had been modified by interaction with lower crustal gabbro. To attribute a carbon enrichment signature to the mantle, we must identify melt inclusions that have avoided  $CO_2/ITE$  ratio modification by crustal interaction.

We have modelled the mixing of a depleted Miðfell melt composition, which has a smooth trace element pattern, with crustal components. We discount a role for significant plagioclase addition to Miðfell melts (Fig. S5). However, mixing with a 10% fractional melt derived from Miðfell gabbro best matches the shape of positive Sr, negative Zr, and Ba > Nb anomalies observed within some



**Fig. 4.** Variation in  $CO_2$  content as a function of Ba concentration for (a) Midfell and (b) Kistufell melt inclusions (diamonds, with vapour bubbles; circles, without vapour bubbles). The maximum  $CO_2$  recovered from melt inclusion glass (circles and unfilled diamonds) is 1170 ppm, as shown by the horizontal line. Combined glass and vapour bubble  $CO_2$  measurements plot between this line and  $CO_2/Ba = 396$  (the maximum ratio preserved by a smooth trace element pattern melt inclusion), as shown by the dashed line. Midfell contains melt inclusions with low enough Ba concentrations that they have likely avoided degassing, while Kistufell melt inclusions are too enriched and have all likely lost  $CO_2$ . One sigma error ellipse plotted in grey.



**Fig. 5.**  $CO_2$  content against (a) Nb abundance and (b) Ba abundance, and  $CO_2/ITE$  against ITE for (c) Nb and (d) Ba, for melt inclusion and glass suites. Miðfell melt inclusions (red) were measured at Edinburgh (circles) and Woods Hole Oceanic Institution (diamonds). Filled diamonds show reconstructed  $CO_2$  content, while empty diamonds are glass only  $CO_2$ . Previously analysed suites are from Siquieros (light blue; Saal et al., 2002), Borgarhraun (royal blue; Hauri et al., 2018), and Equatorial Atlantic (dark blue; Le Voyer et al., 2017) melt inclusions, and MORB glasses (white, Michael and Graham, 2015; sky blue, D-MORB, Shimizu et al., 2016). Dashed lines show constanto:  $CO_2/ITE$  ratios in (a) & (b). Light red regions in (c) & (d) indicate the maximum  $CO_2/ITE$  ratios recorded (from high to low) in: (i) the whole population, (ii) melt inclusions with a smooth trace element pattern, and (iii) melt inclusions show MORB source estimates. One sigma error ellipses are dependent on trace element concentration.

of the Miðfell melt inclusion population (Fig. 3 & S6), providing good evidence for interaction between primitive Miðfell melts and gabbro in the crust.

Extending our analysis to assess the impact of crustal interaction on  $CO_2/ITE$  ratios; an important observation is that carbonate phases have not been seen in Miðfell gabbroic xenoliths (Gurenko and Sobolev, 2006), though carbonate breakdown on decompression during eruption may remove visible evidence of gabbroic carbon (e.g., Canil, 1990). Therefore, the addition of carbon to Miðfell melts cannot be ruled out from textural observations, but geochemical observations can be used to avoid melt inclusions that have strongly interacted with gabbroic material, i.e., those having large trace element anomalies.

To avoid the effects of any potential  $CO_2$ -ITE modification, in the subsequent analysis we have only used melt inclusions with smooth trace element patterns. We believe that these melt inclusions have had minimal interaction with crustal gabbro during ascent through the Miðfell magmatic system.

Melt inclusions with smooth trace element patterns record maximum  $CO_2/Nb = 1832 \pm 316$  and  $CO_2/Ba = 396 \pm 48$  (Fig. 5c,d), again showing higher ratios than observed in MORB suites, and suggesting that the relative carbon enrichment in Miðfell melts likely originates in its source mantle.

# 6. Estimating the carbon content of the Midfell mantle source

The key question we address in this section is whether the observed high  $CO_2/Ba$  and  $CO_2/Nb$  ratios in Miðfell inclusions require an anomalously carbon-rich mantle source. First we identify which ITE's have stayed coupled to carbon during the melting process, and hence which  $CO_2/ITE$  ratios are unfractionated from their source values. We then review the mantle components contributing to Miðfell and calculate in what relative proportions they are represented in its chemistry. Finally, we use constraints on ITE concentrations in these sources to convert the observed  $CO_2$  and ITE systematics of Miðfell inclusions into source  $CO_2$  concentrations.

# 6.1. Carbon - ITE coupling in Miðfell

The silicate-melt partition coefficient for carbon, as estimated by carbonated-lherzolite melting experiments, places it between Ba and Nb in terms of compatibility (Rosenthal et al., 2015). Therefore, we would expect  $CO_2/Ba$  and  $CO_2/Nb$  in Miðfell to be similar to that of its mantle source, provided it represents an aggregation of melts from across the melting region (e.g., Fig. 6 solid line). If instead, Miðfell was produced from high-degree melts from the shallow part of the melting region, a region already extensively depleted by prior melting, then even highly incompatible elements may have been fractionated from each other, so  $CO_2/Ba$ and  $CO_2/Nb$  may not be faithful to the source value (e.g., Fig. 6 dashed and dashed-dotted lines).

We have two tests for whether  $CO_2/ITE$  ratios have been fractionated during melting. The first uses the fact that carbon partitioning between silicate and melt lies between that of Ba and Nb (Rosenthal et al., 2015). Therefore, if Miðfell represents preferential sampling of high-degree shallow melts, we might expect to observe anomalously high  $CO_2/Ba$ . However, this observation would be associated with correspondingly low  $CO_2/Nb$ . Instead, Miðfell inclusions show correlated high  $CO_2/Ba$  and  $CO_2/Ba$  values (Fig. 6), indicating that no such fractionation has occurred.

Secondly, fractionation between trace elements during fractional melting will be manifest in the elements' relative variability (their standard deviation divided by their mean concentration). One prediction of fractional melting models is that trace element variability should increase with decreasing partition coefficient (red line, Fig. 7; Schiano et al., 1993). Fig. 7 shows that in Miðfell inclusions, the most incompatible elements measured (Ba, K, Nb, La) have constant relative variability, which is consistent with a residual porosity during melting having damped the variability generated by low degree melting. As these low degree



**Fig. 6.** The relationship between CO<sub>2</sub>/Nb and CO<sub>2</sub>/Ba in Miðfell melt inclusions. The CO<sub>2</sub>/ITE ratios of the depleted mantle component (DM), as inferred from the Borgarhraun eruption, are shown as grey bars. These ratios represent the starting source composition for modelling fractional melting. The composition of instantaneous melts of this source follow the dashed line, accumulated melts the solid line, and the residual solid the dashed-dotted line. Points along the instantaneous melts curve are labelled by melt fraction. The highest CO<sub>2</sub>/ITE melts cannot be produced by fractional melting of a DM source. Partition coefficients for melting were taken from Rosenthal et al. (2015):  $D_{\rm C} = 0.00055$ ,  $D_{\rm Nb} = 0.0034$ , and  $D_{\rm Ba} = 0.00012$ .



**Fig. 7.** Trace element variability in Miðfell melt inclusions shown by observed relative standard deviation (RSD) of trace elements with varying incompatibility on mantle melting. The results of a pure fractional melting model and the partition coefficients for garnet peridotite melting are shown as a red solid line. The expected behaviour of carbon can be inferred from the blue vertical line, based on its measured compatibility (Rosenthal et al., 2015). Ba, K, Nb, La show similar RSD, suggesting that they remained coupled during melting.

melts will have contained almost all the Ba, K, Nb, and La that was in the source, these elements will not have been fractionated from each other during the melting event that produced Miðfell. As carbon's partition coefficient lies within the range of Ba to La, we can infer from Fig. 7 that carbon will not have been fractionated from any elements across this range of partition coefficients. Therefore, the uncommonly high CO<sub>2</sub>/Ba and CO<sub>2</sub>/Nb ratios observed in undegassed Miðfell melts (Fig. 5) reflect the composition of their mantle sources.

# 6.2. Components in the Miðfell mantle source

*Miðfell's depleted mantle component*: the Borgarhraun eruption provides an on-Iceland sample of the local depleted mantle component in the plume (Thirlwall et al., 2004; Stracke et al., 2003), one that has been suggested to be ubiquitous in the Atlantic mantle (Hauri et al., 2018). Similarities in lithophile elements (Fig. 3), including Pb isotopes (Kokfelt et al., 2006; Halldórsson et al., 2016a), between Miðfell and Borgarhraun, suggest that DMM-like Borgarhraun source material is also present as a component within the Miðfell mantle source.

Is there a pyroxenitic component in Miðfell?: The variability in trace element enrichment and major element chemistry of primitive melts from across Iceland has been interpreted to reflect recycled pyroxenitic components in the Icelandic mantle (Sobolev et al., 2008; Peate et al., 2010; Shorttle and Maclennan, 2011). In the case of Miðfell though, trace element ratios (e.g., Nb/Zr ~0.05), major elements (FeO ~9.4 wt%), and Cl isotopes (Halldórsson et al., 2016a) suggest a minimal contribution from pyroxenite material to this eruption.

This is not to say that Miðfell has had no recycled contribution to its composition. It contains a typical DMM component, which has itself been argued to contain recycled signatures, both as a regional contamination of the upper mantle (Hauri et al., 2018) and as a global phenomenon (Andersen et al., 2015). However, as Borgarhraun does not show significant carbon enrichment with respect to MORB and significantly less relative carbon enrichment than Miðfell, there is no evidence that the Icelandic mantle has been enriched in carbon from pyroxenite addition.

*Evidence for a deep mantle component:* the Miðfell source incorporates material distinct from that of the MORB source, having a more solar- or carbonaceous chondrite-like composition that is often ascribed to an ancient primordial reservoir. This signal is most distinct in the heavy noble gas isotopes (e.g., Harrison et al., 1999; Trieloff and Kunz, 2005; Mukhopadhyay, 2012; Caracausi et al., 2016).

However, the heavy noble gases also indicate that the Miðfell source has had atmosphere recycled into it (e.g., Harrison et al., 1999). This conclusion is in general consistent with constraints from other isotopic systems, e.g., N and Os, which suggest that ancient (> 1.5 Ga) recycled crust is present within the deep mantle component of the Icelandic mantle plume as crustal isotopic signatures are coupled with high-<sup>3</sup>He/<sup>4</sup>He values (Brandon et al., 2007; Halldórsson et al., 2016b). Therefore, a deep mantle component could contain carbon of both primordial and recycled origins.

The above observations suggest that the Midfell source mantle comprises a mixture of a depleted component (DM), akin to Borgarhraun source mantle, and a deep mantle component (Deep). Both the Deep and DM components within the Miðfell source can have plausible estimates made of their ITE content, which will be important for estimating source CO<sub>2</sub> concentrations. In subsequent modelling we adopt the BSE composition given by McDonough and Sun (1995) for the Deep component, though we note that its ITE composition could be more enriched due to recycled material. As there are no constraints on the nature of this enrichment, we do not propagate this uncertainty, although we emphasise that by assuming a BSE source rather than BSE + recycled source, our source carbon estimates will be a lower bound. We assign the DM component the Borgarhraun source mantle ITE composition (Hauri et al., 2018). As the CO<sub>2</sub> of DM has been estimated, only Deep CO<sub>2</sub> and the DM:Deep proportions are unknowns in estimating source CO2. Neither choice of source composition is critical to the results we subsequently obtain: similar source carbon concentrations are inferred if we take, for example, ITE abundances from Palme and O'Neill (2014) for the Deep estimate and Workman and Hart (2005) for the DM.

# 6.3. Proportions of depleted and deep components in the Miðfell mantle source

If Ba, K, Nb, La, and carbon have not been fractionated from each other during melting, then their proportions in Miðfell reflect the degree of melting, F; melt mixing; and melt transport: partial melting will have enriched these ITE's in the melt compared to their source concentrations. Using estimates of Ba, K, Nb, and La content of both DM and Deep components constituting the Miðfell source, we can calculate how much the melting process has enriched Miðfell compared to its mantle source, and thereby calculate the source carbon content. We define the enrichment factor for an element, i, as

$$E_i = \frac{C_i^i}{C_i^0},\tag{1}$$

where  $C_i^l$  is the concentration of the element in the liquid (i.e., observed Miðfell) and  $C_i^0$  is the concentration of the element in the source (DM and Deep taken from Hauri et al., 2018 and Mc-Donough and Sun, 1995, respectively). For Ba, K, Nb, and La we can calculate  $E_i$  directly, assuming a DM:Deep ratio. For carbon, we take the mean enrichment factor estimated from the lithophile trace elements, and use it in (1) with the observed carbon content of Miðfell inclusions to estimate  $C_C^0$ . The main question is what mass fraction of the Midfell source is the Deep component? The enrichment factor contains information on this fraction: given that Ba, K, Nb, and La have not been fractionated during melting, and if we have identified the correct Deep and DM source compositions, then  $E_{Ba} = E_K = E_{Nb} = E_{La}$ . Therefore, by sweeping through Deep component fractions from 0 to 1 and identifying the minimum amount of variability in  $E_i$ , we can identify the optimal source mixture.

In Fig. 8a we show the mean enrichment factor,  $\bar{E}$ , calculated from Ba, K, Nb, and La content of the Miðfell matrix glass (i.e., the composition of the aggregated melt) as a function of Deep fraction in the source. Fig. 8 shows that  $\bar{E}$  lies between 1 and 3 in these models. These small enrichment factors likely reflect both the high peak mantle melt extent under Iceland (~30%; Maclennan et al., 2001) and the incomplete mixing of mantle melts leading to a bias towards relatively shallow fractional melts from the full melting column in the mean Miðfell composition.

When the source is mostly DM (i.e., low  $X_{Deep}$ ) the  $C_i^0$  values are low, and the  $E_i$ 's are correspondingly high. The variability in the calculated enrichment factors is minimised at  $X_{Deep} = 0.47$  (Fig. 8b), i.e., a nearly 50:50 mixture of DM and Deep components in the Miðfell source, which corresponds to  $\bar{E} = 1.56$ .

# 6.4. Translating enrichment factors to source CO<sub>2</sub>

The final step in using the calculated enrichment factors to estimate bulk Miðfell source carbon is to choose a carbon content for the Miðfell magma. For this calculation we take the CO<sub>2</sub> content of the melt inclusions most chemically similar to the matrix glass, which have  $CO_2 = 1079$  ppm: these inclusions likely trap the mixed magma prior to its extensive low pressure degassing, albeit they do not have the highest  $CO_2/ITE$  ratios we observe and therefore may have already lost some  $CO_2$ . Most importantly, this choice is consistent with our use of the matrix glass trace element composition to calculate the enrichment factors. Our calculation is also independent of chosen DM  $CO_2$  value, as it is determined by erupted  $CO_2$  content and  $\bar{E}$  values.

Taking this value of CO<sub>2</sub>, Fig. 8c reports Miðfell source carbon content calculated using enrichment factors over a range of source Deep fractions (blue line). For  $X_{Deep} = 0.47$ , the bulk Miðfell source contains 690 ppm CO<sub>2</sub> (Fig. 8c green bar).



**Fig. 8.** Estimating the fraction of deep mantle component ( $X_{Deep}$ ) in the Miðfell source (a & b), and the CO<sub>2</sub> concentration of the bulk Miðfell source and its depleted (DM) and deep (Deep) components (c). The best-fit mean enrichment factor ( $\tilde{E}$ ) calculated from Ba, K, Nb and La in Miðfell relative to a mixed source (a;  $2\sigma$  error as grey field) is defined by the  $X_{Deep}$  composition where relative standard deviation of the enrichment factor for the four elements is minimised (b; green band). In (c), the Miðfell source CO<sub>2</sub> content was calculated using  $\tilde{E}$  and measured CO<sub>2</sub> (= 1079 ppm). The CO<sub>2</sub> concentration of the Deep component (orange) was calculated by balancing the Miðfell source content with the DM component (105 ± 57 ppm; black; Ba and Nb estimate average from Hauri et al., 2018). 90% confidence envelopes in grey.

Mass balance between the DM and Deep components allows us to convert the bulk source CO<sub>2</sub> (blue) into Deep CO<sub>2</sub> (orange). Here our calculation does depend on knowing the DM CO<sub>2</sub> content, which has been previously constrained as  $105 \pm 57$  ppm (black; Hauri et al., 2018). For X<sub>Deep</sub> = 0.47, Deep has a CO<sub>2</sub> content of  $1350 \pm 350$  ppm (90% confidence interval considering only propagated analytical and  $\bar{E}$  uncertainties).

These calculations are robust for a number of different assumptions: (i) as Fig. 8c shows, the full range of source DM:Deep proportions predict a Deep CO<sub>2</sub> content > 1000 ppm; (ii) if the modelling is repeated using more depleted melt inclusions, which are less likely to have undergone degassing and have peak CO<sub>2</sub> of 300 ppm, then Deep compositions of > 1250 ppm CO<sub>2</sub> are inferred; and (iii) adaptations to the model to explore the consequences of mixing fractional melts from two sources produce the same requirement of > 1000 ppm CO<sub>2</sub> in the Deep source.

#### 7. Summary & discussion

We have shown that melt inclusions from the Icelandic Miðfell eruption record some of the highest CO<sub>2</sub>/ITE ratios reported in basalts, and that these inclusions reflect the composition of their mantle source. In comparison with Borgarhraun, Miðfell's noble gas and lithophile element composition argues for its source comprising a deep mantle component in addition to the depleted mantle common to Icelandic magmas. By combining existing work with our new carbon and trace element observations, we have been able to assign a carbon content to this deep component. We took the depleted mantle component to be that of the well-studied Borgarhraun eruption, which lacks primitive noble gas isotope signatures and has a source  $CO_2 = 105 \pm 57$  ppm (using Ba, Nb estimates; Hauri et al., 2018). We estimate the deep mantle component to have a CO<sub>2</sub> concentration of at least 1000 ppm (preferred value  $1350 \pm 350$  ppm), which increases when the proportion of deep mantle component assumed to be present in the source is decreased.

#### 7.1. Recycled carbon?

Xenon isotopes indicate that a significant proportion of Xe in Miðfell is derived from recycled air ( $\sim$ 90%; Mukhopadhyay, 2012). If carbon was coupled to Xe during recycling, then some amount of carbon in the Miðfell source could also have come from recycling. However, it would be surprising if this component was the origin of the high CO<sub>2</sub>/ITE ratios we observe. Miðfell lavas are not unique in containing recycled air: MORB have been interpreted to have a similar abundance of recycled air in their DMM source (Mukhopadhyay, 2012; Parai and Mukhopadhyay, 2015). Moreover, DMM, in contrast to the Icelandic mantle, is also known from Pb isotopes, U isotopes, and CO<sub>2</sub>/Ba ratios to have been pervasively contaminated by surface material (Andersen et al., 2015; Hirschmann, 2018). Despite this, MORB have lower CO<sub>2</sub>/ITE and inferred source CO<sub>2</sub> concentrations than Miðfell (e.g., Le Voyer et al., 2017). Therefore recycling, at least as seen by MORB, does not appear to significantly enrich carbon with respect to ITE's.

# 7.2. Ancient mantle carbon?

If Miðfell's carbon enrichment is a signature of entrained deep mantle material, to what extent is it ancient mantle carbon? This question is difficult to answer and will require more eruptions to be identified with primitive noble gas signatures and carbon undersaturated melt inclusions. However, the two eruptions we now have on Iceland with CO<sub>2</sub>/ITE ratios that can be linked to their mantle sources, differ almost exclusively in their geochemistry by the presence of primitive noble gas signatures in Miðfell, which are absent in Borgarhraun - their lithophile radiogenic isotope compositions are otherwise very similar (Fig. 1b, c). This signature suggests an association between the primitive noble gases, which have been attributed to solar nebula ingassing during Earth's earliest history (Mukhopadhyay, 2012; Williams and Mukhopadhyay, 2018), and carbon. If true, it would imply a significant fraction of Earth's carbon came directly from the solar nebula rather than later accretion.

However, the coupling of high-<sup>3</sup>He/<sup>4</sup>He with recycled material signatures (Brandon et al., 2007; Halldórsson et al., 2016b) could also suggest that deep mantle carbon is of a recycled origin, making it difficult to differentiate between primordial and recycled carbon from deep mantle material. Especially as there is no certainty that the nature of deep recycled material is the same as the upper mantle recycled material assumed to be present in the Borgarhraun source, which evidences no enrichment in carbon (Hauri et al., 2018).



**Fig. 9.** Estimates of deep mantle carbon reservoir mass (ACO = Atmosphere + Crust + Oceans =  $3.1 \times 10^{23}$  g CO<sub>2</sub>), using the Deep concentration curve in Fig. 8c for a range of mantle fractions (grey dotted lines). Four mantle fractions have been highlighted: (i) 5% representing the seismically-defined D" layer (blue; Tolstikhin and Hofmann, 2005); (ii) 20% for the abyssal layer of the lower mantle, defined by U/K mass balance (green; Arevalo et al., 2009); (iii) 42% for the mass of primitive mantle calculated by <sup>40</sup>Ar mass balance (orange; 90% confidence envelope in grey; this study and Allègre et al., 1996); and (iv) 75% for the lower mantle as defined by the 660 km seismic discontinuity (red; Hofmann, 1997).

#### 7.3. Size of the PM carbon reservoir

If we assume that the Deep carbon estimate derived from Miðfell is representative of the global deep mantle reservoir, then we can calculate a revised mantle carbon budget. Estimates for the deep mantle fraction of the total mantle range from the seismically-defined 660 km transition zone, down to the seismically-defined D" layer above the core-mantle boundary. Fig. 9 presents carbon content estimates of four potential deep mantle reservoir fractions: (i) 5% representing the D" layer (blue; Tolstikhin and Hofmann, 2005); (ii) 20% for the abyssal layer of the lower mantle, defined by U/K mass balance (green; Arevalo et al., 2009); (iii) 42% for the mass of primitive mantle calculated by <sup>40</sup>Ar mass balance (orange; 90% confidence envelope in grey; this study and Allègre et al., 1996); and (iv) 75% for the lower mantle as defined by the 660 km seismic discontinuity (red; Hofmann, 1997).

The carbon content of the atmosphere, crust, and oceans combined (ACO) is  $\sim 3.1 \times 10^{23}$  g CO<sub>2</sub> (Hirschmann, 2018). Carbon stored in the deep mantle, depending on mantle fraction, is therefore up to 14 times greater than the ACO carbon reservoir, while the DMM is approximately equal to the ACO (Fig. 9). Our calculations do not include the potential carbon contribution from the lithospheric mantle, which could be host to one AOC of carbon (Sleep, 2009; Kelemen and Manning, 2015). These carbon reservoir estimates for the deep Earth are necessarily speculative, but comparable to the range of previous bulk mantle estimates (e.g., Halliday, 2013). Our carbon estimates are also a lower bound in two important respects: (i) if the deep mantle component in Midfell is less than the high value we used, then the implied carbon content in it is higher; (ii) if the deep component contains recycled material that is enriched in Ba and Nb, then our use of a BSE composition will have led to an underestimate of its carbon content.

#### **Declaration of Competing Interest**

The authors declare that they have no competing financial interests.

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# Appendix A. Supplementary material

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.epsl.2019.07.002.

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