Rapid core formation in terrestrial planets by percolative flow: in-situ imaging of metallic melt under high pressure/temperature conditions

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Core formation has left a lasting geochemical signature on the Earth. In order to constrain the composition of the Earth we must fully understand the processes by which newly formed Earth, and the bodies which accreted to it, differentiated. Percolation of iron-rich melt through solid silicate has been invoked as a mechanism for differentiation and core formation in terrestrial bodies in the early solar system. However, to date the contribution of percolation to core formation cannot be assessed due to the absence of data on Fe-rich melt migration velocities. Here we use a novel experimental design to investigate textural changes in an analog system, Au melt in polycrystalline h-BN, at 3 GPa, relevant to core formation in the early solar system. Using a combination of high resolution, in-situ X-ray tomography and fast 2-D radiographic imaging, we obtain the first direct data on melt migration velocities at high PT. Melt migration is highly variable and episodic, driven by variations in differential pressure during melt migration and matrix compaction. Smaller scale melt processes, representing migration of melt along pre-existing melt networks, give comparatively fast velocities of 0.6–60 μm s⁻¹. Ex-situ experiments are used to compare melt networks in analog systems to Fe-rich melt in silicates. Two competing processes for melt migration are percolation of melt along grain boundaries, and hydraulic fracturing induced by melt injection. Typically, both processes are noted in experimental and natural systems, although the relative importance of each mechanism is variable. Using a simple model for melt flow through a porous media, migration velocities determined here account for full differentiation of Earth-sized bodies within 10¹–10³ Myr, for submicron diameter melt bands, or within a few Myr or micron-sized melt bands. This is consistent with rapid timescales inferred from geochemistry for core formation in planetesimals, implying that percolation may have had an important contribution to core differentiation in the Earth.

Keywords: core formation, differentiation, planetesimal, tomography, melt migration, percolation, pressure, terrestrial
INTRODUCTION

Core formation represents one of the most significant events in the formation and evolution of terrestrial bodies, including Earth. These bodies have a characteristic layered structure, consisting of metallic, Fe-rich, cores surrounded by thick shells of silicate. This implies that there must have been a mechanism or mechanisms to efficiently separate iron alloy from silicate in the accreted material which ultimately formed Earth, Venus, Mars, Mercury, and other bodies (Stevenson, 1990). Differentiation would have partitioned elements, either fully or partially, between metallic and silicate components, resulting in a lasting geochemical reservoir (Wood, 2008). As such, determining the exact processes of differentiation and core formation, the range of pressure-temperature and P/T conditions under which they operated, and the timescales for these processes, is a critical step in modeling planetary evolution and in constraining the geochemistry of the rocky planets.

During, or shortly after accretion, undifferentiated material in planetesimals and proto-planets underwent reorganization on timescales of millions to tens of millions of years, separating dense Fe-Ni-S core-forming liquids from less dense silicates (Chambers, 2004; Kruijer et al., 2014). In smaller planetesimals (up to 80 km diameter), the heat source for differentiation was likely decay of short-lived isotopes $^{26}$Al and $^{60}$Fe, which will have primarily resulted in melting Fe-S components, then, at higher temperatures, of silicate material (Hevey and Sanders, 2006; Gupta and Sahijpal, 2010). However, transport of heat-producing $^{26}$Al to the surface of smaller bodies in the first silicate melts, coupled with thermal buffering due to Fe-S melting, may have minimized the extent of silicate melting during differentiation (Wilson and Keil, 2012; Mare et al., 2014; Neumann et al., 2014). In larger bodies, prolonged accretion of large impactors resulted in additional heating and more extensive melting, supporting the existence of silicate magma oceans (Nakajima and Stevenson, 2015). Formation of deep magma oceans is typically used to explain the efficiency of core formation in the early solar system (Stevenson, 1990). However, although the existence of long-lived magma oceans can be hypothesized for the Earth, there is growing realization that the magma ocean model cannot alone explain many aspects of core-formation in the early solar system. High-resolution Fe isotopic studies reveal no evidence for terrestrial extreme pressure/temperature fractionation that would be expected in the deep magma ocean model (Craddock et al., 2013; Halliday, 2013). Parent bodies for many meteorites, considered as representatives of bodies which accreted to the proto-planets, show geochemical evidence for low-temperature differentiation in the absence of silicate melting (Mare et al., 2014; Barrat et al., 2015). Furthermore, Rudge et al. (2010) have demonstrated that the observed geochemistry of the Earth can be modeled without the assumption of full metal-silicate chemical equilibration, indicating that core formation in the early solar system system may well have been a protracted, multi-stage process, involving both percolation-aided segregation and later silicate melting. Assessing the limiting conditions under which percolation is a viable mechanism for segregation of Fe-rich melts could help place constraints on the nature and timing of core formation in the inner solar system.

Whilst segregation of Fe-rich liquids from a silicate magma ocean would be a highly efficient mechanism for segregation, the effectiveness of percolation as a mechanism for core formation remains uncertain. Fe-S liquids typically have high dihedral angles in silicate matrices, implying that percolation is only possible above a critical melt fraction (Von Bargen and Waff, 1986; Ballhaus and Ellis, 1996; Shannon and Agee, 1996; Terakawa et al., 2005, 2007). In-situ electrical conductivity studies indicate a connectivity threshold of around 5 vol% (Yoshino et al., 2003, 2004), although other studies reveal that this value might be substantially higher (Walde et al., 2007; Bagdassarov et al., 2009a). A high connectivity threshold for core-forming melts suggests that undifferentiated planetesimals would need a large volume fraction of melt to begin differentiating. However, recent modeling work (Ghanbarzadeh et al., 2017) suggests that in complex natural systems, once interconnected metallic melt networks form, percolation can drain areas of melt to melt fractions of below 2%.

Experimental studies have demonstrated that percolation can act as a mechanism for core formation under certain deep planetary conditions. Numerous studies have highlighted the key role of shear deformation in mobilizing Fe-S rich melts that would otherwise be trapped within solid silicate matrices (Bruhn et al., 2000; Rushmer et al., 2005; Groebner and Kohlstedt, 2006; Hustoft and Kohlstedt, 2006; Rushmer and Petford, 2011). Although Walde et al. (2011) argued that this mechanism of deformation-aided melt segregation does not operate at lower strain rates, the recent study of Berg et al. (2017) demonstrated the efficiency of this mechanism over a wide-range of experimental strain rates under high PT. Shi et al. (2013) demonstrated that extreme pressure also plays a fundamental role in modifying textures in equilibrated Fe-S liquid-silicate systems, implying that percolative flow would have occurred in the lower mantles of the early Earth and Venus. Terakawa et al. (2008) have also shown that full interconnectivity of Fe-S liquids can occur under more oxidizing conditions regardless of deformation, which might explain low-temperature differentiation in planetesimals. Therefore, percolative core formation can be initiated in systems with high metallic fractions, and in systems with low metallic fractions under oxidizing conditions, under extreme pressures or during deformation. Once initiated, percolation can effectively drain core forming melts from a system. In fact, the fraction of trapped Fe-rich melt left within the silicate matrix is small enough to satisfy observed highly siderophile element concentrations in the bulk silicate Earth (Ghanbarzadeh et al., 2017).

Geochemical constraints imply timescales for planetary core formation of, at most, tens of millions of years (Kleine et al., 2009), and as little as 3 Myr after formation of Ca-Al-rich inclusions in planetesimals (Kleine and Wadhwa, 2017). Therefore, it remains uncertain whether Fe-rich liquid segregation is rapid enough for percolation to have played a significant role. Most studies of core-forming melt percolation rely on textures to indirectly infer migration velocities, and are typically based on gravity driven percolation through a stable melt network (e.g., Hustoft and Kohlstedt, 2006; Roberts et al., 2007; Bagdassarov et al., 2009b; Kohlstedt and Holtzman, 2009).
As such, melt migration velocities are, at best, guestimates. Recent studies (Bagdassarov et al., 2009b; Cerantola et al., 2015) have also shown that partial melting of silicate matrices can actually inhibit segregation of Fe-rich melts, implying that if percolation is a viable mechanism for core formation, it must be efficient enough to result in Fe-melt segregation before the onset of silicate melting. Bagdassarov et al. (2009b) provide one of the only direct experimental assessments of Fe-melt segregation velocities in silicate, based on a high-pressure/temperature centrifugal study. Using textural analysis of quenched samples they modeled segregation of FeS melt within a compacting silicate matrix, and derived segregation velocities of $10^{-4}$ to $10^{-5}$ mm/h, scaled to 1 g gravity. These rates would be too slow to efficiently form planetary cores, which would require minimum segregation velocities of $10^{-3}$ mm/h for Earth, or $10^{-2}$ mm/h for Vesta. However, Bagdassarov et al. (2009b) derived melt segregation velocities indirectly from 2-D textural analysis of run products, which provide limited understanding of progressive textural development in dynamically evolving samples. Recently, Todd et al. (2016) provided the first assessment of Fe-melt migration velocities based on in-situ measurements in analog core-segregation experiments. They obtained high resolution 3-D X-ray tomographic images of a sample of 4.5% FeS melt in polycrystalline olivine during progressive annealing at high PT and during torsional deformation. However, the method which they used provided a limited voxel size of 2 µm; previous experimental studies have demonstrated that melt networks in deforming olivine-FeS systems typically develop at the micron scale or smaller (Berg et al., 2017), and only scale to larger features at very high strain rates when extensive melt channeling develops. Todd et al. (2016) determined melt migration velocities indirectly using Lattice-Boltzmann simulations, which were then used to simulate flow through meshes created from their 3-D textural datasets. However, importantly, the image thresholding technique used by Todd et al. (2016) was based partly on expected melt contents of radiographic images; as such, at the resolution available, finer melt networks were not resolved, and permeability may have been significantly underestimated. Furthermore, they did not observe melt textures previously described in olivine-FeS high pressure deformation experiments (Walte et al., 2011; Berg et al., 2017), suggesting that deformation-aided melt segregation was not simulated in the experiments.

Recently, Philippe et al. (2016) described a modified Paris-Edinburgh type apparatus (RoToPEC) for performing in-situ tomographic studies at extreme conditions (high pressure/temperature/stress), similar to the previously described Paris-Edinburgh type deformation apparatus of Bromiley et al. (2009) used in the core-formation experimental study of Berg et al. (2017). Philippe et al. (2016) additionally provided initial in-situ synchrotron X-ray tomography data on evolution of Fe-S melt rich regions in polycrystalline olivine, under various high pressure/temperature conditions during torsional deformation. As with the study of Todd et al. (2016), voxel resolution was limited to around 2 µm, and Philippe et al. (2016) were also unable to resolve fine core-forming melt channels, or assess any melt migration in their in-situ tomographic data set. In this study, we develop an alternative approach to determine melt migration velocities under extreme conditions. This approach is based on initial in-situ studies described above, but in a simplified analog system, and with a modified experimental setup aimed to produce melt channels of a sufficient size to be resolved within the current limitations of high-pressure and/or tomographic synchrotron beamline facilities.

**METHODS**

**In-Situ Tomography**

The RoToPEC apparatus was installed on the ID27 extreme conditions beamline at the European Synchrotron Radiation Facility (ESRF), Grenoble, France, using a custom-built stage capable of moving in X, Y, and Z orthogonal directions to align the sample within the X-ray beam. A motorized Soller slit system was used to collected in-situ diffraction data, coupled with PerkinElmer image plate (Mezouar et al., 2002). Radiographic images were obtained using a MARCCD 165 camera, with the automated system allowing rapid toggling between diffraction and imaging modes. X-ray camera control was coupled to the motors driving rotation of the RoToPEC anvils, allowing projections to be taken at increments of 0.02° during 180° sample rotation. Exposure time and beam energy were varied based on attenuation of the sample volume and surrounding gasket during trial experiments, with optimized values of 0.37 Å beam wavelength, equivalent to $\approx 33$ keV, and an 100 ms exposure time.

**Figure 1** shows the sample assembly. A boron-epoxy gasket was used with an internal graphite resistance furnace, molybdenum rings to ensure good electrical contact with the upper and lower anvil faces, and alumina spacers. A split-capsule design was used to promote melt migration through the central portion of the assembly visible during in-situ radiography. This consisted of upper and lower hexagonal boron nitride (h-BN) polycrystalline capsules with a central, drilled volume housing the melt-rich assemblage. The intention of this design was to promote considerable melt migration and channelization along the central join between the 2 sides of the capsule. An outer single piece h-BN sleeve was used to enhance stability of the sample volume. h-BN was used in this study as the matrix material, after a series of unsuccessful trial experiments, for a number of reasons: (1) the availability of high quality, fine-grained polycrystalline material with a minimal variation in grain size, (2) the ease with which the material could be machined accurately, and (3) its higher thermal conductivity, which was observed to minimize blowouts during sample melting, presumably due to lower thermal gradients during the initial stages of experiments. Although h-BN differs in key characteristics from mantle silicates, as we discuss later, gold melt in h-BN provides useful insight in metallic melt mobility under mantle conditions. Gold was used as the melt phase as it is a commonly used analog material for investigating textural development in silicate/core-forming melt studies (e.g., Bruhn et al., 2000; Walte et al., 2011). Although the melting point of gold is several hundred degrees lower than Fe-S, at high-pressures, this facilitates long duration experiments. Central regions of the sample were loaded with a mix of 20 vol% fine-grained
FIGURE 1 | RoToPEC sample assembly. (A) Schematic showing sample assembly, consisting of a shaped boron-epoxy gasket containing integral graphite resistance furnace. (B) 3-D depiction of central sample volume used in melt migration study, consisting of a 2-piece polycrystalline h-BN capsule containing drilled central volume (gray) housing 2-phase mixture of polycrystalline gold and powdered h-BN. (C) 2-D schematic of sample volume shown in (B) with additional outer boron nitride sleeve to aid sample stability at high PT. Gray transparent region represents portions of the sample assembly visible to the X-ray beam, which is otherwise absorbed by the upper and lower carbide anvils.

(0.5–0.8 μm) spec-pure gold powder mixed with powdered h-BN (<1 μm), homogenized in a agate pestle and mortar by grinding in air for 30 min. Pressure and temperature were determined using existing calibrations for hydraulic load (oil pressure) and power (Bromiley et al., 2009; Philippe et al., 2016; Berg et al., 2017), and from diffraction data using known equations of state for both gold and h-BN (Le Godec et al., 2000; Shim et al., 2002). Diffraction patterns were also used to determine onset of gold melting, from the rapid disappearance of gold reflections in diffraction patterns. Figure 2 shows a schematic of how data was collected during the experiment once run pressure had been obtained.

Ex-Situ Experiments
Additional experiments were conducted at the University of Edinburgh to investigate processes for Au melt migration in h-BN and MgO, and their relevance to Fe-S melt migration in polycrystalline silicate. A series of end-loaded piston-cylinder experiments were performed with using the same capsule design, for capsules made of polycrystalline h-BN and MgO. Run conditions, heating and cooling cycles replicated those of the RoToPEC experiment, with additional comparison experiments performed using a single heating/pressurization stage, such that the sample was fully pressurized, heated to above the Au melting point, and then quenched after a period of 24 h. Full details of the piston cylinder apparatus, sample assembly and calibration are given in Bromiley et al. (2010). All recovered samples were analyzed using a Carl Zeiss SIGMA HD VP Field Emission Scanning Electron Microscope (SEM). Samples proved to be challenging to polish due to the extreme contrast in hardness between fine Au melt networks and h-BN /MgO. However, recovered samples typically fractured longitudinally to the compression axis of compression of the experiments, meaning that sample surfaces were typically quite flat, and could
be easily characterized by secondary electron imaging, and in some cases, by backscattered electron imagining.

In-Situ Experimental Details and Data Processing

After pressurization to 3 GPa over 1 h, a tomography scan at room temperature was taken, and the sample was then heated to 1,000°C at a rate of 50°C/min. Temperature was maintained for an additional 2 h before heating at 10°C/min until melting of Au was observed. Temperature was then dropped to below the melting point of Au and a series of images taken. Over the next 20 h, pressure was continually monitored and maintained whilst temperature was cycled through the melting point of Au. Changes in Au melt distribution were too fast to be tracked by tomographic scans, which took 45–60 min to acquire. Therefore, a combination of high-spatial resolution 3D tomographic scans and high temporal resolution 2-D radiographic movies were used to monitor and characterize textural changes. A further 3 melting events were monitored during which (1) 3-D tomographic scans were obtained while the sample was held just below the melting point of Au; (2) 2-D radiography movies obtained while the sample was held above the melting point of Au for a few tens of minutes; (3) a 3-D tomographic scan obtained after the temperature had been rapidly lowered to solidify the Au melt. Voxel size in tomography scans was 2.8 µm, although the actual resolution was slightly lower than this. Reconstruction of 2D slices from the tomographic dataset was performed using the University of Ghent’s Octopus 8.7 reconstruction software (Dierick et al., 2004) using a filtered background projection algorithm to create sonograms and 2D slices from projections. Flat field and dark current images were obtained before and after each tomographic scan by driving the sample out of the beam path, and then closing the experimental shutter. Octopus software was used to remove ring artifacts during the creation of sinograms, minimize the effects of beam hardening and apply a simple noise filter during reconstruction. 2D projections had an exposure time of 100 ms, a time interval of 20 s between projections, and a spatial resolution of 2.5 µm. Projections for each movie were normalized using flat field and dark current images to improve sample contrast and enhance signal-noise ratios. To highlight changes between projections as a function of time, an initial reference frame, p₀, was subtracted from each subsequent frame, pₓ, to create a “difference” image, pₓ–p₀, using the ImageJ Image Calculator function (Abramoff et al., 2004). Grayscale in resulting images represents change in sample attenuation between the reference and new projection.

RESULTS FROM IN-SITU TOMOGRAPHY AND RADIOGRAPHY

Figure 3 shows volume renderings from tomographic datasets. Scan 1, obtained before heating, shows that some misalignment between the 2 halves of the capsule occurred during loading/pressurization. This misalignment is obvious in all subsequent scans, although there does not appear to be any significant slip between the capsule halves during the experiment (i.e., subsequent to initial pressurization and annealing). In
fact, high resolution scans (for example Figures 8, 9) show that there is a single, coherent melt-rich region in the central sample volume. Furthermore, we note no evidence for an influence of this misalignment on melt network development, with some melt sheets cutting across the original join between the capsule halves. As discussed below, however, it is likely that melt distribution in the sample influences textural development.

In addition to misalignment, there is also increased beam attenuation at the join between the 2 halves of the capsule prior to melting, mostly likely representing Au grains dropped onto the h-BN capsule during loading of the central melt-rich volume, and/or minor amounts of extrusion of the softer Au along the capsule join during pressurization. From subsequent volume renderings it is clear that (1) considerable Au melt migration throughout the h-BN outer capsule occurred during the heating cycles, (2) that no observable melt extrusion occurred along the capsule join, and instead that (3) a complex pattern of melt channels/sheets developed in the sample, with both vertical and horizontal movement of Au melt through the h-BN. It is also evident that (4) a fraction of the Au melt escaped the h-BN capsule during the experiment, typically through vertical to near vertical melt channels. During each heating cycle there was further shortening of the sample volume (the height of rendered volumes shown in Figure 3 is constrained by the spacing between carbide anvils). As is typical in Paris-Edinburgh cell experiments, progressive annealing of samples at elevated P-T results in sample shortening, and the effect here was markedly increased during melt migration.

The melt network which forms within the sample consists predominantly of extensive, thin melt sheets (Figure 3; see also higher magnification images in Figures 8, 9). These are sub-planar to undulate, typically on the order of 10 µm thick, 10–100 µm wide, and extending 100–1,000 s µm in length. Less extensive more arcuate melt sheets are also noted. Melt sheets are generally anastomosing, frequently bifurcate and interconnect. Once formed, melt networks typically persist throughout numerous heating cycles, although some features, especially those adjacent to the edges of the sample volume, disappear during progressive heating, implying melt drainage from regions of the sample. There is evidence for extensive horizontal and vertical movement of melt, with other sheets forming at oblique angles to the axis of compression. Locally, at terminations of sheets there is also evidence for melt ponding. Toward the bottom of the sample volume the dominant features are thinner melt bands, typically vertical to near vertical (Figure 4). Limitations in resolution (at best several microns) mean that larger melt channels are easier to detect in in-situ data, and there could be additional, extensive melt migration through much smaller melt features which would not be detected in this dataset. Changes in texture between later heating cycles are much less obvious, and consistent with the slow drainage of melt from the sample, accumulation of melt at the bottom and edges of the sample volume, and loss of melt to other parts of the sample assembly, largely through pre-existing melt channels/sheets.

Radiographic movies demonstrate both larger scale, faster horizontal and vertical melt migration events early in heating cycles (Figure 5), and smaller scale movements later in heating cycles (Figure 6). Figure 5 shows a series of stills collected over the first 160 s of the final heating stage of the experiment. px-p0 images show clear evidence for considerable lateral movement
of melt through an undulating melt network, resulting in rapid accumulation of a large volume of melt toward the outer edge of the h-BN capsule. After this period of rapid melt migration, changes in melt distribution are considerably reduced, as shown in Figure 6. Variations between $p_x - p_0$ images are much reduced, allowing smaller, active melt channels, especially those resulting in localized variation in attenuation, to be readily identified. Two such discrete movements, M2 and M3, are labeled in Figure 6, on images where melt migration was first observed. Melt migration velocities can be determined by tracking melt movement along fractures between projections at fixed time intervals, as shown in Figure 7. Melt velocities are not constant, and even accounting for scatter, show changes in velocity. This mirrors more obvious changes in melt migration velocities observed during larger scale melt migration processes. Figures 8, 9 show examples of how melt migration events can be matched in radiographic projections and tomographic reconstructions to constrain melt migration velocities. Correlation of 2D radiographs and 3D

FIGURE 4 | (A) Whole volume renderings for scans 5, 7, and 8 (shown in Figure 3) oriented to show a basal perspective of the sample volume. Au melt shown in gold, and h-BN capsule in transparent white for reference, mm-wide, $10^1 \mu m$ thick melt sheets with a vertical to near vertical orientation are evident. (B) Corresponding 2D reconstructions from close to the base of the sample volume, showing both permeating melt sheets (fuzzy white) and linear fracture-like sheets. Numbers on vertical axis refer to scan numbers. There is a significant change between scans 5 and 7 due to extensive melt accumulation at the base of the capsule, and some melt loss out of the sample volume. Several melt features evident in 5 are absent in 7, presumably as melt has drained away. In contrast, there is minimal evidence for additional fracturing between scans 7 and 8, as discussed in the main text.
FIGURE 5 | Radiographic projections highlighting Au melt movement during the final heating stage of the experiment. Vertical axis shows total time in s. Left: original projects, Right: projection minus initial projection to highlight changes in the sample volume. Black regions represent increases in attenuation due to accumulation of Au melt, most noticeable in the meandering horizontal melt channel and melt “pocket” toward the bottom the sample volume. There is a corresponding decrease in attenuation in the central melt area due to outward migration of Au melt. After 160 s (to final image) no further significant changes were noted in the sample for a further 5 min.

...tomographic reconstructions works best for these smaller scale movements due to complexities in melt networks. The absence of changes in fracture geometry between scans obtained before and after this heating cycle also implies that these movements utilized pre-existing melt flow channels, or planes of weakness. As such, extracted velocities constrain inherent timescales of melt migration and percolation, as opposed to timescales of melt sheet developed through, for example, some hydraulic fracturing process.

For melt migration events, complexities in the melt structure meant that a number of possible flow paths could be identified which might represent the true melt path observed in 2D radiographic images. In each case, maximum, minimum, and most likely melt displacements were determined. These estimates are shown in Table 1. Velocities determined in 3 well-constrained, distinct events, 0.6–60 μ m.s⁻¹ (or 0.1–5 m.hr⁻¹) are comparable to those in other small scale melt events throughout the sample which are more difficult to accurately constrain. Estimated velocities from events such as the protracted, large scale horizontal redistribution of melt shown in Figure 5 are at least one order of magnitude faster. Therefore, velocities vary by 2–3 orders of magnitude.

METALLIC MELT MIGRATION PROCESSES IN VARIOUS SYSTEMS

As h-BN contrasts markedly from silicate materials which constitute the interiors of rocky planets such as Earth, it is important to demonstrate that data obtained here is of use in constraining core-forming processes in the inner solar system. Optical and SEM images of run products (Figure 10) can be used to compare melt networks in Au melt-h-BN and Au melt-MgO systems to Fe-rich melt in silicate.
The *in-situ* and h-BN piston-cylinder experiments showed comparable textures (Figures 10A,B). h-BN used in these experiments was very fine-grained (∼1 µm diameter equant grains). Au melt characteristically formed extensive, undulating, but approximately planar melt sheets around 10 µm thick, but up to 100 s µm wide. Melt sheets were horizontal to sub-vertical, and allowed melt to readily migrate to the edge of the capsule and pool against graphite furnaces. In melt-rich regions of samples there was additional formation of melt networks resulting from extensive wetting of h-BN grain boundaries. Successive heating cycles resulted in more extensive melt migration away from the capsule center (original melt-rich region), although planar melt networks were well-developed in all samples.
Experiments performed using MgO capsules produced different textures. Optical examination revealed that Au melt had permeated extensive volumes of the MgO capsules, and even Al₂O₃ spacers placed adjacent to the capsule, resulting in an obvious gold/brown discoloration (Figures 10C,F). SEM images demonstrate that this is due to: (1) extensive grain boundary wetting and formation of meandering melt networks, and (2) Au melt trapped in grain boundary regions representing melt-drained regions of the sample, i.e., residual Au melt (Figures 10D,E). Au melt moved horizontally into the surrounding MgO, but also vertically. Melt sheets were noted in regions of very high melt fraction (Figures 10H,I), with sheets tending to be sub-horizontal. Additional experiments were performed varying Au melt fraction in the capsule center from 1:3 to 3:1. At high melt fractions, melt sheets were more likely to develop, comparable to sheets seen in h-BN.

In both MgO and Al₂O₃ there is a clear influence of matrix grain size on texture (Figure 11). Au melt pockets size scales with matrix grain size, implying that grain boundary wetting is the main process for melt migration. In addition, the fraction of trapped melt in fine grained regions is higher due to the higher volume fraction of grain boundaries. Melt trapped in finer grained regions is typically present as nm-sized spherical “blobs” in contrast to the well-defined melt pockets present in coarser grained regions (Figure 11). This might indicate that Au melt more readily migrated through finer-grained material, an interpretation supported by visual examination of recovered run products. The low volume fraction of Au in melt drained regions of samples is consistent with Ghanbarzadeh et al. (2017); as hypothesized, percolative flow can effectively drain melt from the solid matrix, leaving a much lower melt fraction than would be estimated from a critical melt threshold based on dihedral angles.

Melt migration by extensive grain boundary wetting, and brittle-type fracturing resulting in planar-type melt sheets, have both been observed in analog experiments performed in the system FeS melt + polycrystalline olivine (Berg et al., 2017). In that study, torsional deformation was used to constrain the influence of strain on grain boundary wetting at very low melt fractions. This contrasts with the present study where high melt
fractions promoted development of melt networks. Berg et al. (2017) noted that fracturing occurred readily at higher strain rates in the FeS-olivine system, although fractures are limited in extent compared to the Au + h-BN experiments, and extend only a few 10s of μm across a small number of olivine grains. In most of the experiments of Berg et al. (2017), performed at lower strain rates, grain boundary migration dominated. Although there is a clear influence of grain size, it is apparent that brittle fracturing and melt sheet formation is more prevalent in the analog h-BN experiments, and grain boundary wetting more prevalent in other analog materials (Au melt in MgO and Al₂O₃, FeS melt in olivine). Figure 12 highlights the differing
FIGURE 9 | Matching melt migration event M2 in 2D radiographs to 3D tomographic reconstructions. (A) An example of an original radiographic projection (p70), with approximate start and end positions for the observed melt movement labeled (from Figure 6). (B) Image p70-0 highlighting melt movement in the projection. (C) 3D tomographic reconstruction. In this instance it is more difficult to identify the exact melt channel as there are multiple possibilities, i.e., start and end positions. Estimates of the shortest and longest routes are shown. In this case, 2D radiography shows good evidence for melt movement along a pre-existing fracture network. (D) Measurements of the shorter route between start and end locations, with an intermediate, slightly more convoluted route (mid-estimate). (E) Measurements of the longest estimated route from the sample middle (3) to the edge of the field of view.
nature of these two processes. For grain boundary migration, channels connect melt pockets trapped at grain junctions. For all analog core-forming experiments described here, wetting angles of the melt are characteristically high, which means that melt interconnectivity can only proceed at melt fractions above the critical melt threshold (here) or during deformation (Berg et al., 2017). As such, grain boundary regions have a dominant influence on melt migration and textural development. In contrast, hydraulic fracturing results in formation of planar melt sheets which indiscriminately cut across grain boundaries and grain centers.

Experiments demonstrate that: (1) Au melt moves relatively quickly through polycrystalline h-BN even at extreme PT conditions, despite the high interfacial tension and inferred low mobility of Au melt; (2) The dominant process for rapid Au melt migration appears to be hydraulic fracturing and the development of melt sheets, rather than grain boundary migration, especially during the first heating stage on the in-situ (h-BN) experiment, or early stages of later heating cycles; (3) however, in later heating cycles, it is possible to extract data on melt migration rates in pre-existing melt networks; melt migration rates determined here are nevertheless relatively fast, of the order of microns to tens of microns per second; and (4) it is clear that melt migration is episodic, with periods of rapid melt mobility interspersed with longer periods of quiescence, during heating cycles.

### DRIVING FORCES FOR MELT MIGRATION

Melt in the in-situ experiment is observed to move through pre-existing fractures at rates of $\sim 10^{-6}$ to $10^{-5}$ ms$^{-1}$. Gold melt is expected to have low mobility due to its high surface tension and high dihedral angle of 145$^\circ$ in cubic h-BN (Ricci and Novakovic, 2001). Melt movement observed can be modeled using a simple Darcy’s flow calculation for sheet flow along a single planar fracture (i.e., cubic law) of the form:

$$q = \frac{\rho g w^3 bh}{\mu 12 \delta L}$$  \hspace{1cm} (1)

where $q$ = advective velocity, $\rho_i$ = density of melt, $w$ = fracture aperture, $\mu$ = melt viscosity and $\delta h/\delta L$ is the hydraulic gradient. $h$ is the hydraulic head, which can be expressed as a combination of the elevation head ($z$) and the pressure head ($P/\rho g$), defined as $h = z + P/\rho g$. At high $T$ but ambient $P$, $\mu$ for Au melt is estimated at $0.04$ Pa.s (Ofte, 1967), and $\rho_i$ at $17,310$ kgm$^{-3}$. $L$ can be taken as $\approx 0.001$ m, and $z \approx 1$ mm. If flow of Au melt is produced simply as a result of change in elevation (i.e., $P/\rho g = 0$), $q$ is calculated to be of the order of $10^{-12}$ ms$^{-1}$, at least 6 orders of magnitude lower than observed. This calculation ignores factors such as fracture roughness, and fracture tortuosity, which may both reduce flow rates; regardless, it is clear that an additional driving force is required, especially as this mechanism would also not explain horizontal melt movement.

By varying values of $\delta P$, the hypothetical fluid overpressure required to produce melt migration on the order observed can be estimated over a range of feasible aperture values, giving a range of solutions of $0.001$–$5$ MPa. A factor which could act as a driver for melt migration is fluid over-pressurization as a result of a phase change, i.e., melting of the Au during the initial stages of each heating cycle. Assuming that the h-BN matrix is relatively incompressible, the overpressure will be equivalent to the pressure difference, $\delta P$, between the pressure required to compress the liquid from its theoretical density at atmospheric $P_i$, $\rho_l$, to the density of the solid state $\rho_s$, and the background confining pressure, $P_c$:

$$\Delta P = \left( K_T \frac{\Delta \rho}{\rho_l} \right) - P_c$$  \hspace{1cm} (2)

where $K_T$ is the bulk modulus and $\Delta \rho$ the difference in density between solid and liquid states. Assuming $\rho_l$ for Au is $1.93 \times 10^3$ kgm$^{-3}$, $\rho_l$ is $17310$ kgm$^{-3}$ and the bulk modulus of liquid gold is $63$ GPa at $1,600$ K (Steck and Bek, 1976; Blairs and Joasoo, 1980), and taking into account a $3$ GPa pre-existing confining pressure, the overpressure associated with Au melting is $\sim 3$ GPa. This drops to $2$ GPa when considering that gold in the central volume of the capsule is mixed with h-BN in a ratio of $1:3$ (density $2,100$ kgm$^{-3}$), but is still several orders of magnitude higher than that required to cause flow at the velocities observed, for apertures of $1$–$20$ µm.

If overpressure due to melting was the sole cause of melt migration, melt movement in all directions through the outer capsule might be expected. Melt channels in varying orientations are observed, although there appears to be a preference for melt to move sub-vertically at the bottom of the sample volume. This could indicate an additional gradient encouraging melt flow in certain directions. The density difference between h-BN and Au is large ($\sim 15,000$ kgm$^{-3}$), which will create a subtle melt gradient, although the driving force is much lower than that of fluid overpressure. More likely, a thermal gradient may be present within the sample volume, undetermined but probably of the order of tens of K or more; thermal gradients have previously been inferred to result in significant melt movement in other partial melt, high PT experiments (Leshner and Walker, 1988).

Large fluid overpressure in the Au melt-h-BN system explains why brittle fracturing was observed during heating cycles, especially immediately after melting. Conversely, prior to each heating stage, compression of the sample volume is consistent with the pressure required to move the liquid through fracture plumes.
FIGURE 10 | (A). Secondary electron image showing extensive Au melt sheet(s) in h-BN outer capsule. These sheets (100 s µm wide and long, but a few µm thick) acted as conduits for Au melt to leave the capsule center (melt-rich region) and pool at the interface between the h-BN and furnace. (B) Magnified view of same sample, showing that most Au is retained in the capsule center, but that there are meandering melt sheets through the outer h-BN capsule and accumulation of melt adjacent to the graphite furnace (black). (C) Section across center of MgO capsule. Texture is different to experiments with h-BN, with Au melt forming extensive melt networks permeating through the coarser-grained MgO capsule, and far fewer broad melt channels. (D) Back-scatter electron image (BSE) showing Au melt (light gray) in MgO matrix (dark gray) in same sample, showing interconnected Au in the center of the sample (bottom) and an interconnecting melt channel extending upwards. Trapped melt is also present in the surrounding MgO. Au melt loss from the capsule, and accumulation adjacent to the furnace, was also noted. (E) BSE image of part of same sample, showing thin Au melt channel wetting MgO grain boundaries. Grain size in MgO is ~10 µm (compared to <1 µm in h-BN). Pockets of Au melt show typically high dihedral angles in MgO. Submicron blobs of Au melt along grain boundaries represent trapped (residual) melt. (F) Magnified image of Au in MgO experiment with a higher fraction of Au melt in the capsule center (75%). Extensive staining of the surrounding MgO capsule is evident, along with some Au melt sheets. Au melt migrated horizontally and vertically, including into the alumina spacers above and below the capsule. (G) BSE image of same sample showing Au melt (white) in MgO, forming extensive sheets at the boundary between the inner and outer capsule. Higher magnification image (H) shows that brown coloration in (F) is due to submicron blobs of residual Au trapped at grain boundaries. These represent trapped residual melt in dried up melt networks, which mainly transported melt vertically through the sample volume. (I) Similar texture showing extensive Au melt sheet with high dihedral angle in polycrystalline Al2O3 in BSE image of the bottom spacer adjacent to the MgO capsule; once again, fine blobs of Au are interpreted to be residual melt.

with plastic deformation, as is gradual shortening of the sample throughout the experiment as melt migrates away from the inner capsule. Brittle processes occurring at high PT conditions have been observed in other experimental studies, both from observed microstructures (Dellangelo and Tullis, 1988) and from in-situ acoustic measurements (Dobson et al., 2002). Observations in real rock exposures show ductile fabrics, implying high PT conditions, overprinted with evidence of contemporaneous brittle fracturing by melt. These can be explained by the process of "melt-enhanced embrittlement" (Davidson et al., 1994), again linked to overpressure created by melting. Brittle processes are thought to occur when fluid/melt overpressure is higher than can be relieved by percolative flow along grain boundaries. As such, the process is more common in low permeability systems, and would be expected to occur here where the melt is a large volume of high surface-tension liquid, where flow along grain boundaries is limited, and where significant overpressures can build up. Differences between textures in h-BN (grain size ≈ 1 µm or smaller) and MgO/Al2O3 (grain size ≈ 10 µm) suggest an influence of grain size on textural development. Within MgO and Al2O3 there is a significant control of grain size on melt textures, with increased melt trapping (i.e., less drainage of melt) and significantly reduced thickness of melt channels/sheets in fine-grained regions. Thinner melt bands in finer grained h-BN could inhibit percolative flow, and enhance hydraulic fracturing.
The majority of laboratory and field observations made during several heating cycles. This implies that overpressure of secondary melt migration events may be significantly lower than the original hydraulic fracturing event, leading to lower fluid flow velocities, consistent with observations made during several heating cycles. This implies that the slower movement seen during in-situ experiments provides a meaningful assessment of melt migration velocities. As such, data on Au melt migration velocities in h-BN determined here are independent of the original mechanism for promoting development of melt networks, fast melt injection of melt associated with over-pressurization during melting. The key point here is that we constrain migration velocities through pre-existing melt networks, independent of the mechanism for initial melt network development and channelization.

**MELT MIGRATION VELOCITIES IN EXPERIMENTAL AND NATURAL SYSTEMS**

In contrast to experiments performed here, Philippe et al. (2016) failed to note any evidence for Fe-S melt migration in polycrystalline olivine. In that study, a significantly lower melt...
End-member mechanisms for migration of metallic melt in polycrystalline material under extreme P-T conditions. (A) Formation of melt channels (red) by grain boundary wetting (grains pried apart; dominant influence of grain boundaries). (B) Melt migration by fracture propagation (grains fractured, with little influence of grain boundaries). Panels (C,D) show formation of melt channels/sheets by both processes in the system FeS melt-olivine (Berg et al., 2017) in deformation experiments at extreme P-T. In these experiments, fracturing becomes dominant at higher strain rates, while grain boundary migration dominates at low strain rates, although fractures in the olivine matrix are typically a few 10's of µm in length at most and cut across a small number of olivine grains, as opposed to the (typically) extensive sheets noted in Au + h-BN experiments. Panels (E,F) show comparable features in the present tomographic data set (Au melt in h-BN) observed in tomographic scans. More diffuse, poorly defined, regions of increased attenuation are interpreted as extensive wetting of grain boundaries, while hydraulic fracturing produces well-defined, planar features.
fraction was used, which could explain the lack of a driving force, such as overpressure, for melt migration. Furthermore, although volume expansion of Fe-S on melting is higher than Au, the higher compressibility of Fe-S, as a volatile-bearing melt, means that potential overpressures during the phase change are lower. Using Equation (2), assuming a central melt-rich zone of 20% FeS + 80% olivine, and using data for FeS from Urakawa et al. (2001), the pressure difference upon FeS melting at around 2 GPa is approximately −1 GPa. Even if the center of the sample was 100% melt, the FeS melt overpressure would only be 0.1 GPa, an order of magnitude lower than in Au melt experiments. As such, the only driving force for melt migration outwards in the experiments of Philippe et al. (2016) is non-hydrostatic stress and a small thermal gradient. Rushmer et al. (2005) estimated that shear deformation in partially molten systems, at strain rates of \(10^{-5}\) to \(10^{-8}\) s\(^{-1}\), will produce differential stresses within individual melt pockets on the order of 0.1–100 MPa, several orders of magnitude lower than overpressures produced here by melting of Au. As such, differential stresses are unlikely to cause fracturing observed in Au melt experiments. However, melt overpressures arising from deformation may still act to cause fluid flow into grain boundaries. Assuming differential stress is the only driving force for melt migration, velocities calculated using (1) for flow along planar sheets into grain boundaries of around 0.1 µm (consistent with observations in Berg et al., 2017), are on the order of \(10^{-11}\) to \(10^{-9}\) ms\(^{-1}\). This corresponds to melt migration distances of the order of 0.1–10 µm for an hour long heating cycle. Whilst this is consistent with observations from Berg et al. (2017), who described melt channels of 10s–100s µm in length for heating/deformation experiments lasting several days, time constraints for in-situ experiments, and resolutions in tomographic scans of the order of µm mean that comparable experiments designed to determine migration velocities of Fe-S melt may be challenging to perform. Furthermore, unless experiments were designed with a large pressure/temperature/stress gradient, there is no clear driving force to encourage significant melt migration.

An additional factor affecting melt migration, and textural development in partially molten, core differentiation analog experiments, is the physical dimensions of melt channels. Melt channels/sheets formed by hydraulic fracture during the in-situ experiments are typically a 1–10 µm in diameter. Flow rate, according to (1), is proportional to the cube of aperture size, which allows us to assess the effect of melt textures on measured flow rates. Melt channels/sheets/networks which develop in other analogs systems, such as in polycrystalline MgO and Al\(_2\)O\(_3\), or during deformation of Fe-S + olivine, are typically an order of magnitude smaller. This implies flow rates through these finer melt networks around 3 orders of magnitude slower than in Au melt- h-BN experiments, or \(6 \times 10^{-4}\) to \(6 \times 10^{-2}\) µm\(^3\)s\(^{-1}\). For a 1 h heating cycle, this would imply melt movement of distances between ~2 and 200 µm. This comparison assumes that melt migration velocities for Au melt and Fe-S, within different matrices, are comparable. However, melt migration distances of 2–200 µm are consistent with visual observations of longer duration ex-situ experiments performed here, in which Au melt migrated outside of the entire sample volume, and observations of Fe-S migration in longer annealing/deformation studies (Berg et al., 2017), where submicron melt bands permeate 100s of µm after annealing times of several days. This latter observation is important, as it demonstrates that melt migration velocities obtained here, for Au melt in h-BN, are comparable with studies in more realistic analog core-formation systems, such as Fe-S in olivine, validating the analog system used in this study. It also demonstrates that scaling given in (1) is sufficient to extrapolate data to different systems, where mechanisms for melt network development differ. Scaling the range of melt migration velocities (2–200 µm h\(^{-1}\)) observed for FeS melt in olivine in submicron to micron-wide melt sheets to planetary scales implies that core-forming melts can migrate the radius of the Earth over timescales of 40 Myr to 4 Gyr. This lower end estimate is within the 10s Myr constraint for core formation within the Earth. However, core-forming melt segregation from solid silicate through micron-sized melt networks is likely only the initial stage of differentiation through percolation. Progressive percolation will result in increased channelization of melts, and formation of melt-rich bands (e.g., Hustoft and Kohlstedt, 2006). These 1–10 µm melt bands would permit crust to core melt migration in 4 Myr, even given the lowest measured melt migration velocity determined here. Alternatively, Fe-S melt migration velocities calculated above from (1), assuming differential stress as a driver for melt movement through an olivine matrix, can be used to independently estimate timescales for core segregation. For submicron melt channels, this method predicts core formation via percolation of the order of 2 Gyr timescales. However, this reduces to 2 Myr for micron sized melt channels, consistent with timescales estimated from Au melt analog experiments. As such, metallic melt migration velocities determined here are consistent with a significant role for percolation, prior to silicate melting, in early core-formation and planetary segregation in rocky planets and smaller bodies.

Similar textures to those noted in Au and Fe melt percolation experiments have been observed in achondrite meteorites, and used to infer melt flow in the interiors of planetesimals. McCoy et al. (2006) noted both Fe infilled fractures (presumably developed due to over-pressuring or internal deformation) and pervasive Fe wetting of silicate grain boundaries in melt-poor regions of the meteorite GRA 95209. Chen et al. (2002) noted extensive Fe-S melt injection into shock fractures in the Sixiangkou meteorite, but also presented textural evidence for melt filled fractures and wetting of silicate grain boundaries in regions adjacent to shock veins. As such, both textural and geochemical (Mare et al., 2014; Barrat et al., 2015) evidence from meteorites suggests that percolation contributes to redistribution of Fe-rich melts in parent bodies (large asteroids or planetesimals). Experimental studies demonstrate that percolation is a viable mechanism for Fe-melt segregation, and therefore, core segregation, under various conditions (Bruhn et al., 2000; Terasaki et al., 2008; Shi et al., 2013; Berg et al., 2017). Modeling work further demonstrates that once initiated, percolation can efficiently drain Fe-melt from solid silicate to low melt fractions. Results here now suggest that percolation can result in rapid Fe-melt segregation, consistent with the timescales for core formation in the early solar system. Work is required
to determine exact mechanisms for Fe-S melt percolation in silicates, how these vary as a function of composition, pressure, temperature and stress, and corresponding melt migration velocities. However, it is clear that the contribution of percolation to core formation in Earth and other terrestrial bodies needs to be considered. The assumption of one-stage, full core-mantle chemical equilibration at the base of a terrestrial magma ocean is overly simplistic; more likely, core formation was a multi-stage, protracted process which will have left a more complex geochemical signature on terrestrial reservoirs.

AUTHOR CONTRIBUTIONS

MB conducted most of the data analysis presented as part of her Ph.D. project. GB supervised the project, devised the experimental programme with MB, and conducted ex-situ experiments. YL assisted with in-situ experiments and helped design and machine the sample assemblies. MM, J-PP and JP both assisted with the in-situ experiment and initial interpretation of results during the experiment, and MM led modifications of the ID27 beamline to allow installation of the RoToPEC press and integration with beamline software. NP conducted SEM analysis of recovered run products. GB and MB wrote this manuscript, and other authors contributed to preparation of the final draft.

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