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Published in:
Geochemistry, Geophysics, Geosystems

Link to article, DOI:
10.1002/2013GC005047

Publication date:
2014

Document Version
Publisher's PDF, also known as Version of record

Citation (APA):
Almeida, T. P., Muxworthy, A. R., Williams, W., Kasama, T., & Dunin-Borkowski, R. E. (2014). Magnetic characterization of synthetic titanomagnetites: Quantifying the recording fidelity of ideal synthetic analogs. Geochemistry, Geophysics, Geosystems, 15(1), 161-175. https://doi.org/10.1002/2013GC005047

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Magnetic characterization of synthetic titanomagnetites: Quantifying the recording fidelity of ideal synthetic analogs

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[1] A series of four synthetic basalts comprising titanomagnetite (Fe$_{3-x}$Ti$_x$O$_4$) grains of varied size and titanium content have been produced by a glass-ceramic method. Complementary characterization techniques of X-ray diffractometry, secondary electron microscopy, and transmission electron microscopy (TEM) demonstrate the reaction product composition consisted of mainly Fe$_{3-x}$Ti$_x$O$_4$, pyroxene hedenbergite, fayalite, and SiO$_2$. The samples exhibit bimodal distributions of larger (<2 μm) and smaller Fe$_{3-x}$Ti$_x$O$_4$ particles (<50 nm in diameter), the latter found inside pyroxene crystals, as well as the sporadic occurrence of dendritic Fe$_{3-x}$Ti$_x$O$_4$ structures. Magnetic measurements show their bulk characteristics fall into two groups: Ti-rich titanomagnetite samples with varying Ti content; and near-stoichiometric magnetite. The TEM technique of off-axis electron holography allowed for visualization of the magnetic behavior of the synthetic Fe$_{3-x}$Ti$_x$O$_4$ grains. Energy dispersive X-ray analysis and off-axis electron holography confirmed the small Fe$_{3-x}$Ti$_x$O$_4$ grains (<50 nm) confined within glassy pyroxene regions to be Fe-rich and single domain, carrying strong magnetic signals, compared to the relatively magnetically weak larger Fe$_{3-x}$Ti$_x$O$_4$ grains (x ~ 0.6). The large grains in the pure magnetite sample are shown to be pseudo-single domain in nature. The quenching process involved in synthesis is considered similar to that of pillow basalts found at mid-ocean ridges and hence the reaction products are thought ideal in terms of characterization and understanding, for the purpose of studying natural systems.

Components: 5,582 words, 15 figures, 2 tables.

Keywords: titanomagnetites; palaeomagnetism; electron holography.

Index Terms: 1540 Rock and mineral magnetism: Geomagnetism and Paleomagnetism; 1599 General or miscellaneous: Geomagnetism and Paleomagnetism; 1021 Composition of the oceanic crust: Geochemistry.

Received 16 September 2013; Revised 4 November 2013; Accepted 6 November 2013; Published 27 January 2014.
1. Introduction

[2] Our knowledge of the geomagnetic field’s variance on geological timescales is dependent on our ability to recover meaningful palaeomagnetic signals from rocks and sediments. Hence, it is important to understand the formation and recording mechanisms, as well as fidelity, of the magnetic minerals within natural systems. The most common magnetic minerals on Earth, and the main recorders in igneous rocks including mid-ocean ridge basalts (MORB), fall within the iron-titanium-oxide system [Dunlop and Özdemir, 1997] and generally provides the only recoverable absolute palaeointensities (as well as palaeo-directions). Such full vector data is essential to fully reconstruct the ancient geomagnetic field behavior.

[3] Although the iron-titanium-oxide system has been widely studied over the years, there are still gaps in our knowledge. Most previous studies have looked at natural systems [e.g., Bleil and Petersen, 1983; Krása et al., 2005], with only a few investigations focusing on synthetic samples where the magnetic minerals formation history is tightly constrained [e.g., Özdemir and Banerjee, 1981; Halgedahl, 1991; Zitzelsberger and Schmidbauer, 1996]. Key questions still remain as to the formation of Ti-rich iron oxide minerals in the common single domain (SD) to pseudo-single domain (PSD) grain-size range: (1) what are the micromagnetic structures within Ti-rich iron oxide PSD grains, how do these affect their recording fidelity and (2) what is the variation in stoichiometry of the particles. Evidence suggests there is a stoichiometric variance with grain size [Zhou et al., 1997, 2000] which will give rise to misleading Curie temperatures if determined by an in-field thermomagnetic measurement (an $M_s$-$T$ curve, where $M_s$ is the saturation magnetization and $T$ is the temperature), i.e., the smallest grains carrying the remanence will have a different stoichiometry to the larger grains that dominate the $M_s$-$T$ curve.

[4] In this paper, a microscopic and magnetic study of a suite of iron-titanium-oxides synthesized in the laboratory using the glass-ceramic method developed by Worm and Markett [1987], is reported. This technique has been refined to gain a better control of the grain size and spatial distribution, allowing the synthesis of Ti-rich iron oxides in the SD to PSD grain size range. The transmission electron microscopy (TEM) technique of off-axis electron-holography [Dunlop-Borkowski et al., 1998; Harrison et al., 2002; Kasama et al., 2013] has been applied to synthetic Ti-rich PSD iron oxides for the first time, and an investigation of the behavior of SD and PSD grains as a function of temperature is presented.

2. Methods and Experimental Procedures

[5] Synthetic basalts containing Fe$_{3-x}$Ti$_x$O$_4$ were prepared by a glass-ceramic method, as described previously [Worm and Markert, 1987]. Samples were synthesized from laboratory grade powders of Fe$_2$O$_3$, SiO$_2$, TiO$_3$, CaCO$_3$, K$_2$CO$_3$, and Na$_2$CO$_3$, as summarized in Table 1. The powders were mixed in acetone using a pestle and mortar and once dry combined with aqueous polyethylene oxide solution to form a viscous slurry and pressed into pellets (~5 mm in diameter) on the end of Pt wires. The pellets were suspended in a vertical temperature controlled tube furnace under an oxygen partial pressure provided by a CO-CO$_2$ gas flow of ratio 1:10. The sample was melted at 1400°C for 15 h to equilibrate Fe$^{2+}$/Fe$^{3+}$ ratios with the oxygen fugacity. The Pt wires were then released and the molten pellets subsequently quenched in the bottom of the tube furnace. After quenching the pellets were reheated at 750°C for 3 h using the same method, and quenched again.

[6] For the purpose of structural characterization, quenched pellets were crushed into a fine powder using the pestle and mortar, dispersed on a Si substrate and examined using X-ray diffractometry (XRD) (PANalytical X’Pert PRO Diffractometer). Additional pellets were mounted in epoxy resin and polished for the purpose of survey secondary electron microscopy (SEM) (FEI Quanta 200 ESEM FEG), whilst their elemental constituents...
were identified using energy dispersive X-ray analysis (EDX) (Aztec, Oxford Instruments). Desirable regions of the polished samples were isolated, ion-milled into lamallae and lifted out using a focused ion beam (FIB) and Omniprobe system (Helios NanoLab 600) for localized examination using TEM. Conventional bright field (BF) imaging was performed using a FEI Tecnai TEM operated at 200 kV (Centre for Electron Nanoscopy (CEN), Technical University of Denmark), with EDX analysis (Aztec, Oxford Instruments) and selected area electron diffraction (SAED) allowing for chemical and phase identification, respectively. For the purpose of electron holography, a FEI Titan with a Cs corrector on the condenser lens, equipped with a Lorentz lens and biprism, was used at 300 kV (CEN). Off-axis electron holograms were recorded in Lorentz mode with the objective lens switched off and the specimen in magnetic-field-free conditions at room temperature. Mean inner potential contributions to the recorded phase shift were removed by using in situ magnetization reversal [Kasama et al., 2011].

The magnetic measurements were conducted at the palaeomagnetic laboratory at Imperial College London and the Institute for Rock Magnetism (I.R.M), University of Minnesota. High-temperature thermomagnetic curves were measured using a Princeton Measurements Vibrating Sample Magnetometer (VSM) fitted with a furnace; heating was performed in flowing He. Low-temperature hysteresis measurements were made from 15 to 295 K (in 10 K intervals) using a Princeton Measurements VSM. A Quantum Designs Magnetic Properties Measuring Systems (MPMS) cryogenic magnetometer was used to measure cooling and warming curves of room temperature (RT) induced saturation isothermal remanence magnetization (SIRM) to identify low-temperature crystallographic transitions, as well as warming curves of SIRM at 10 K after field cooling (FC) at 2.5 T and zero-field cooling (ZFC), from 300 K to 10 K.

3. Results

3.1 X-Ray Diffraction analysis

The XRD patterns of Figure 1 provide information on the crystal structure of both the synthesized Fe$_{3-x}$Ti$_x$O$_4$ particles and supporting glass-ceramic matrix. The peaks in Samples S1–S4 are in good agreement with the presence of Fe$_{3-x}$Ti$_x$O$_4$, and characteristic to the phases Fe$_2$TiO$_4$ (Joint Committee on Powder Diffraction standards (JCPDS) reference 72-1665), Fe$_{2.5}$Ti$_{0.5}$O$_4$ (JCPDS 75-1376), and Fe$_2$O$_4$ (JCPDS 88-0315). In addition, all XRD patterns display peaks characteristic to the pyroxene hedenbergite (CaFeSi$_2$O$_6$, JCPDS 70-1876), andradite (Ca$_3$Fe$_2$(SiO$_4$)$_3$, JCPDS 84-1938), FeSi (JCPDS 86-0795), SiO$_2$ (JCPDS 83-0540), and Fayalite (Fe$_2$SiO$_4$, Figure 1. XRD patterns of Samples S1–S4 confirming the presence of Fe$_{3-x}$Ti$_x$O$_4$, indexed to Fe$_{2.5}$Ti$_{0.5}$O$_4$ (JCPDS 75-1376), as well as the additional phases of CaFeSi$_2$O$_6$, Ca$_3$Fe$_2$(SiO$_4$)$_3$, FeSi, and SiO$_2$. 

Table 1. Summary of Precursor Powders Blended into Synthetic Melt Compositions

| Precursor   | Sample S1 | Melt (%) | Sample S2 | Melt (%) | Sample S3 | Melt (%) | Sample S4 | Melt (%) |
|-------------|-----------|----------|-----------|----------|-----------|----------|-----------|----------|
| SiO$_2$     | 0.287     | 28.7     | 0.287     | 30.1     | 0.431     | 34.8     | 0.287     | 31.4     |
| CaCO$_3$    | 0.155     | 15.5     | 0.155     | 16.2     | 0.233     | 18.8     | 0.155     | 17.0     |
| Na$_2$CO$_3$| 0.061     | 6.1      | 0.061     | 6.4      | 0.092     | 7.4      | 0.061     | 6.7      |
| K$_2$CO$_3$ | 0.061     | 6.1      | 0.061     | 6.4      | 0.092     | 7.4      | 0.061     | 6.7      |
| Fe$_2$O$_3$ | 0.349     | 34.9     | 0.348     | 36.4     | 0.348     | 28.1     | 0.348     | 38.2     |
| TiO$_2$     | 0.087     | 8.7      | 0.043     | 4.5      | 0.044     | 3.5      | 0       | 0        |
| **Total**   | **1**     | **100**  | **0.955** | **100**  | **1.240** | **100**  | **0.912** | **100**  |
JCPDS 71-1673), with the latter more dominant in Sample S4.

### 3.2. Scanning-Electron Microscopy Imaging and Analysis

The backscattered-electron (BSE) SEM images (Figure 2) provide information on the size and morphology of the Fe$_{3-x}$Ti$_x$O$_4$ particles dispersed in the glass-ceramic matrix. In samples S1 and S2 (Figures 2a and 2b respectively), the Fe$_{3-x}$Ti$_x$O$_4$ particles (bright regions) are generally isotropic and $\sim 1.6 \mu$m in size, with clusters of smaller Fe$_{3-x}$Ti$_x$O$_4$ particles in the latter. Figure 2c shows sample S3 to comprise slightly larger Fe$_{3-x}$Ti$_x$O$_4$ particles ($\sim 2.8 \mu$m), whilst sample S4 is observed to consist of markedly smaller Fe$_{3-x}$Ti$_x$O$_4$ particles ($\sim 0.7 \mu$m, Figure 3d). Allocation of an appropriate gray value threshold for the BSE SEM images allowed for creation of binary contrast images, with the extrapolated particle-size distribution and nearest-neighbor distance data of samples S1–S4 summarized in Figure 3 [Beggan and Hamilton, 2010]. However, the particle size analysis is limited by the resolution of the SEM and resulting pixel size, with sizes below $<100$ nm considered unreliable for the magnification of the acquired images.

The BSE SEM and corresponding EDX compositional maps (Figure 4) present the elemental distribution within sample S1. Comparison of the BSE image (Figure 4a) with the Fe and Ti compositional maps (Figures 4b and 4c) confirm the bright regions in the BSE image are rich in Fe and Ti, and hence attributed as Fe$_{3-x}$Ti$_x$O$_4$ particles. The remaining area is shown to be rich in Si (Figure 4d) and assigned as the glass silicon matrix, which exhibits areas high in Ca content (Figure 4e), considered as hedenbergite phase regions. Figure 4f shows the oxygen content to be uniformly distributed across the sample.

![Figure 2. (a–d) BSE SEM images of Samples S1–S4, where Fe$_{3-x}$Ti$_x$O$_4$ particles (bright regions) are dispersed within the glass-ceramic matrix.](image-url)
3.3. Transmission Electron Microscopy Imaging and Analysis

[11] The BF TEM images of Figure 5 provide a closer examination of the Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} particles and surrounding glass ceramic matrix. Figure 5a reveals the various sizes of the Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grains in Sample S1, as confirmed by SAED (Figure 5a, inset, indexed to Fe\textsubscript{2.5}Ti\textsubscript{0.5}O\textsubscript{4} (JCPDS 75-1376)). A large single crystal Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grain in the PSD region (>500 nm) is presented in Figure 5b, as confirmed by SAED (Figure 5b, inset). Figure 5c shows a much smaller Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grain, surrounded by the Ca-rich hedenbergite phase, as identified by characteristic lattice fringes corresponding to \{220\} planes (Figure 5c, inset). The associated EDX spectra of Figure 5d, acquired from the Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grains presented in Figures 5b and 5c, show a markedly weaker Ti signal acquired from the smaller Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grain (Figure 5c). The BF TEM image of Sample S4 (Figure 5e) displays large Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grains (>500 nm), distinct from smaller Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grains (<100 nm) which appear to be concentrated in the Ca-rich glassy hedenbergite regions of the matrix, as identified in the SEM data. Examples of the larger Fe\textsubscript{3-x}Ti\textsubscript{x}O\textsubscript{4} grains (>300 nm) in Sample S4 are shown in Figure 5f.

[12] Figure 6a presents a high angle annular dark field (HAADF) image of Sample S4, showing larger Fe\textsubscript{3}O\textsubscript{4} grains and Ca-rich glassy regions confining smaller Fe\textsubscript{3}O\textsubscript{4} grains (<50 nm). The EDX maps of Figures 6b–6d acquired from the boxed region in Figure 6a depicts the (b) Fe, (c) Si, and (d) Ca content. The bottom right corner of the boxed region (arrowed in Figures 6a, 6b, and 6d) of the matrix is absent of both Fe and Ca, further suggesting the glassy regions within the SiO\textsubscript{2} matrix are CaFeSi\textsubscript{2}O\textsubscript{6} or Ca\textsubscript{3}Fe\textsubscript{2}(SiO\textsubscript{4})\textsubscript{3}, whilst the smaller Fe-rich Fe\textsubscript{3}O\textsubscript{4} grains are clear to see. The SEM and TEM images of samples S1 (Figure 7a) and S4 (Figure 7b), respectively, display the dendritic structures found sporadically across these synthetic basalts. Dendritic structures are normally associated with rapid quenching [Dantzig and Rappaz, 2009].

Figure 3. (a–d) Particle size distributions of Samples S1–S4, along with nearest neighbor distance (inset).
The off-axis electron holography presented in Figures 8–10 allowed for the imaging of the in-plane magnetization of Samples S1 and S4, respectively. Figure 8a shows an electron hologram of sample S1 comprising several larger Fe$_{3-x}$Ti$_x$O$_4$ grains (labeled a) and two smaller Fe$_{3-x}$Ti$_x$O$_4$ particles (~50 nm, labeled b). The magnetic induction map of Figure 8b depicts the magnetic field contours to be concentrated to the smaller Fe$_{3-x}$Ti$_x$O$_4$ particles (magnified in Figure 8c), where the magnetic field direction is imaged as downward (Figure 8d) through SD Fe$_{3-x}$Ti$_x$O$_4$ grains. Figure 9a presents an electron hologram of an individual Fe$_{3-x}$Ti$_x$O$_4$ grain (~350 nm) in Sample S4, where the corresponding magnetic induction map shows a magnetic vortex (Figure 9b). The electron hologram of Figure 10a displays an individual Fe$_{3-x}$Ti$_x$O$_4$ grain (~220 nm), again in sample S4, where the corresponding magnetic induction map (Figure 10b) shows its PSD (vortex) nature, which contains both a magnetic vortex and a stray magnetic field similar to that expected for a SD particle.

### 3.4. Magnetic Measurements

High-temperature thermomagnetic analysis indicated that the samples were thermally stable (Figure 11). Curie temperatures were determined using the second-derivative method [Tauxe, 1998] (Table 2). Samples S1 and S3 yielded two distinct Curie temperatures: S1, 83°C and 569°C; and S3, 119°C and 579°C; whereas sample S2 had a broad Curie temperature range, 382–432°C, and a Curie temperature at 579°C; whilst sample S4 exhibited a single Curie temperature of 579°C. The higher Curie temperatures are indicative of near-stoichiometric magnetite, whereas the lower Curie temperatures suggest the presence of the second phase, with significant Ti content.

Measurement of the following room-temperature hysteresis parameters are summarized in Table 2: the remanent saturation magnetization $M_{rs}$; saturation magnetization $M_s$; coercivity of remanence $B_{cr}$; and coercivity $B_c$. All four samples display hysteresis parameters that are traditionally indicative of PSD hysteresis behavior. It is observed that all samples are plotted within the PSD region, however, sample S1 has a high $M_{rs}/M_s$ ratio, which is indicative of SD-like behavior, whilst sample S4 displays the least SD-like characteristics.

First-order reversal curve (FORC) diagrams [Roberts et al., 2000] were measured for the four
samples (Figure 12). All the samples displayed PSD/SD type behavior with distinct peaks [Maxworthy and Dunlop, 2002]. The FORC distribution of Sample S3 is markedly different from the other three, with extension along the $B_c$ axis. Samples S2 and S3 display FORC distributions that slope slightly toward lower $B_u$ values as $B_c$ increases; this is indicative of negative mean-interaction.
fields [Pike et al., 1999]. The near-stoichiometric magnetite sample S4 exhibited behavior characteristic of ideal single-domain grains, however, no negative region close to the vertical $B_\text{cr}$-axis is observed, indicating that the controlling anisotropy is not uniaxial, but likely a mixture of cubic and uniaxial anisotropy [Muxworthy et al., 2004; Newell, 2005].

[17] Low-temperature SIRM warming curves (Figure 13) display a range of behaviors. In all four samples, the sharp drop in the remanence on initial cooling is indicative of very small nanometric grains, which are superparamagnetic at room temperature. Sample S4 (Figure 13d), displays behavior typical of near stoichiometric magnetite with a Verwey transition of $\sim$120 K. The large drop in the remanence at the Verwey transition suggests that the magnetic remanence is dominated by the magnetocrystalline anisotropy. Samples S1, S2, and S3 all display kinks in their SIRM warming curves $\sim$50–80 K, which is indicative of $x = 0.4$–0.6 in $\text{Fe}_3\text{Ti}_x\text{O}_4$ [Moskowitz et al., 1998], though the relatively small drops in magnetization compared to Maskowitz et al. [1998] is attributed to the smaller mean grain size of the samples in this study. Sample S1 also displays a kink in the FC warming curve at $\sim$270 K, which is not seen in the ZFC curve and hence likely due to unblocking of SD grains.

[18] Low-temperature cycling of the SIRM to 10 K (Figure 14), reveals different trends: samples S1 and S2 both demagnetize on cooling without displaying a sharp Verwey transition, sample S4 displays similar behavior, though the Verwey transition is more clearly defined, whilst sample S3 does not display significant demagnetization during cooling; its behavior is almost reversible. The behavior of samples S1 and S2 is typical of titanomagnetite with a low Ti-content with a PSD domain state [Özdemir and Dunlop, 2003], whilst sample S4 is indicative of near-stoichiometric PSD magnetite [Muxworthy and McClelland, 2000]. In contrast, the behavior of S3 is typical for Ti-rich or SD titanomagnetite [Muxworthy and McClelland, 2000].

[19] The hysteresis parameters $B_\text{cr}$ and $M_{rs}/M_s$ are plotted as a function of temperature (15 K to 295 K) in Figure 15. Sample S4 displays behavior for both $B_\text{cr}$ and $M_{rs}/M_s$ typical of PSD/MD magnetite [Muxworthy, 1999], e.g., on warming $B_\text{cr}$ drops
sharply between 100 and 120 K, before increasing as the temperature increases to the room temperature. In contrast, samples S1–S3 display a general decrease in $B_c$ and $M_{rs}/M_s$ on warming from 15 K to 295 K with no pronounced change in behavior around the Verwey transition, which is typical of Ti-rich titanomagnetite [Hodych and Matzka, 2004].

4. Discussion

[20] All four samples were formed by quenching, making their formation history more similar to the pillow basalts found at mid-ocean ridges than the slowly cooled subaerial basalts found elsewhere. The effect of rapid cooling is clearly seen by the presence of dendritic structures (Figure 7) [Shaar and Feinberg, 2013]. In all the samples there appear to be bimodal distributions of particles: larger particles which are readily observed under the SEM data (Figure 2), and smaller particles (<50 nm in diameter), which are found inside pyroxene crystals (Figure 6). These smaller crystals are only seen under the TEM in this instance. It is believed this is the first report of magnetic inclusions within the synthetic pyroxene “single-crystals” [Tarduno and Cottrell, 2005]. Such magnetic inclusions are thought to be ideal recorders of the ancient magnetic field, as they are both

Figure 8. (a) Electron hologram of Sample S1 comprising several larger Fe$_{3-x}$Ti$_x$O$_4$ grains (labeled a) and two smaller Fe$_{3-x}$Ti$_x$O$_4$ particles (~50 nm, labeled b), with the arrows showing direction of the applied saturating field. (b) Corresponding magnetic induction map recorded at remanence after saturating the elements upwards, showing magnetic field contours concentrated to the two smaller Fe$_{3-x}$Ti$_x$O$_4$ grains. (c) Magnification of the two smaller Fe$_{3-x}$Ti$_x$O$_4$ particles (labeled b in Figure 8a). (d) Corresponding magnetic induction map recorded at remanence and arrows show the local magnetization directions. The contour spacing is 0.098 radians and magnetization direction is depicted in the color wheel (inset).
small, and enclosed and protected from weathering. That these grains can be easily grown in the laboratory is encouraging, and shows that such “single-crystals” are not unique, but likely ubiquitous, i.e., the scope for recovering and studying such crystals is high. The larger crystals, which are thought to contain more titanium, display PSD vortex structures similar to those reported for stoichiometric PSD magnetite. In these still relatively small particles, the effects of magnetostriction are not so readily observed in the domain patterns when compared to larger Ti-rich iron oxides [Moshkowitz et al., 1988]. This implies that the magnetic remanence carried by PSD titanomagnetite crystals is not so strongly effected by local stress variations and is a better recorders of the geomagnetic field than the remanence of larger Ti-rich iron-oxide grains.

[21] The bulk characteristics and behavior of the four samples fall into two groups: samples S1, S2, and S3, and sample S4. Samples S1, S2, and S3 are Ti-rich titanomagnetite samples with varying Ti content, whilst sample S4 is indicative of near-stoichiometric magnetite; however, within the titanomagnetite rich samples, there is variability that is not consistent, e.g., S2 and S3 display similar magnetic hysteresis parameters (Table 2), though
samples S1 and S3 display similar thermomagnetic curve behavior (Figure 11) and samples S1 and S2 similar FORC diagrams (Figure 12). The low-temperature Curie values for Samples S1 and S3 are in good agreement with, and hence can be attributed to, grains with the approximate stoichiometry of Fe$_{2.4}$Ti$_{0.6}$O$_4$ [Dunlop and Özdemir, 1997]. Sample S4 consistently displays the behavior expected for PSD/SD magnetite.

The electron holographic imaging also sheds light on some of the other properties. For example, S1, S2, and S3 were observed to have both a range of grains sizes and mixed mineralogy as identified from the thermomagnetic analysis. The electron holographic imaging found that the smaller grains were generally far more magnetic than the larger grains. From this we infer that the smaller grains are more Ti-poor titanomagnetite like, as supported by EDX analysis (Figure 5d), and contribute to the near-stoichiometric magnetite Curie temperatures (Table 2), and the larger grains are Ti-rich titanomagnetites giving rise to the low Curie-temperature phase, i.e., the titaniun preferentially resides in the larger titanomagnetite crystals. The flip side of this statement is that the smaller SD-like better magnetic recorders are more likely to be small SD-like magnetite crystals. The problem remains that commonly the magnetic mineralogy of a sample is determined by measuring the Curie temperature in a relatively large magnetic field: this maybe misleading as the magnetic remanence carriers are not necessarily the biggest contributors to high-field thermomagnetic curves.

**Table 2.** Hysteresis Parameters and Curie Temperatures

| Sample | $B_{r}$ (mT) | $B_{c}$ (mT) | $M_{r}$ / $M_{s}$ | $T_{c}$ (°C) |
|--------|--------------|--------------|------------------|-------------|
| S1     | 14           | 18           | 0.45             | 83 and 569  |
| S2     | 19           | 40           | 0.33             | 382–432 and 579 |
| S3     | 28           | 55           | 0.39             | 119 and 579 |
| S4     | 27           | 58           | 0.24             | 579         |

Figure 11. Thermomagnetic curves of samples (a) S1, (b) S2, (c) S3, and (d) S4. Heating was performed in flowing helium, in a field of 100 mT.
Figure 12. FORC diagrams for samples: (a) S1, (b) S2, (c) S3, and (d) S4. The smoothing factor (SF) for each of the diagrams was two. The measurement time was 250 ms.

Figure 13. SIRM warming curves from 10 K to 300 K for samples (a) S1, (b) S2, (c) S3, and (d) S4, after field-cooling (FC) and zero-field cooling (ZFC) from room-temperature. A field of 2.5 T was applied.
Figure 14. Room-temperature SIRM cycling curves from 300 K to 10 K to 300 K for samples (a) S1, (b) S2, (c) S3, and (d) S4. The SIRM was induced in a field of 2.5 T (check).

Figure 15. Coercive force and $M_r/M_s$ versus temperature for: (a) S1, (b) S2, (c) S3, and (d) S4.
[23] We believe these samples are ideal in terms of characterization and understanding, for studying natural systems under controlled conditions.

5. Conclusions

[24] Synthetic basalts considered similar to pillow basalts found at mid-ocean ridges have been successfully produced by a glass-ceramic method. The reaction products are composed mainly of Fe$_{3-x}$Ti$_x$O$_4$, pyroxene hedenbergite, fayalite, and SiO$_2$. Magnetic measurements show their bulk characteristics fall into two groups: Ti-rich titanomagnetite samples with varying Ti content, and near-stoichiometric magnetite. In the Ti-rich titanomagnetite samples, off-axis electron holography showed the small Fe$_{3-x}$Ti$_x$O$_4$ grains (<50 nm) confined within glassy pyroxene regions to be Fe-rich and single domain, carrying strong magnetic signals, as compared to magnetically weak larger Fe$_{3-x}$Ti$_x$O$_4$ grains (x ~ 0.6). In contrast, the large grains in the pure magnetite sample are shown by off-axis holography to be pseudo-single domain in nature. These reaction products are thought to be ideal in terms of characterization and understanding, for the purpose of studying natural systems.

Acknowledgments

[25] This project was primarily funded by NERC grant NE/H00534X/1, with additional funding from the Royal Society and the Institute for Rock Magnetism, University of Minnesota, USA, which is funded by the W. M. Keck Foundation, The National Science Foundation, and the University of Minnesota.

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