Identification and environmental interpretation of diagenetic and biogenic greigite in sediments: A lesson from the Messinian Black Sea

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Abstract Greigite (Fe3S4) is a widespread authigenic magnetic mineral in anoxic sediments and is also commonly biosynthesized by magnetotactic bacteria in aqueous environments. While the presence of fossilized bacterial magnetite (Fe3O4) has now been widely demonstrated, the preservation of greigite magnetofossils in the geological record is only poorly constrained. Here we investigate Mio-Pliocene sediments of the former Black Sea to test whether we can detect greigite magnetofossils and to unravel potential environmental controls on greigite formation. Our magnetic analyses and transmission electron microscope (TEM) observations indicate the presence of both diagenetic and bacterial greigite, and suggest a potentially widespread preservation of greigite magnetofossils in ancient sediments, which has important implications for assessing the reliability of paleomagnetic records carried by greigite. TEM-based chemical and structural analyses also indicate the common presence of nickel-substituted diagenetic iron sulfide crystals with a ferrimagnetic greigite structure. In addition, our cyclostratigraphic framework allows correlation of magnetic properties of Messinian Black Sea sediments (Taman Peninsula, Russia) to global climate records. Diagenetic greigite enhancements appear to be climatically controlled, with greigite mainly occurring in warm/wet periods. Diagenetic greigite formation can be explained by variations in terrigenous inputs and dissolved pore water sulfate concentrations in different sedimentary environments. Our analysis demonstrates the usefulness of greigite for studying long-term climate variability in anoxic environments.

1. Introduction

Greigite (Fe3S4) is a widespread iron sulfide mineral that has been found in anoxic sedimentary environments across the world over geologically significant time periods [Roberts et al., 2011a]. It is a strongly ferromagnetic mineral that makes important contributions to paleomagnetic and environmental records. Paleomagnetic records carried by greigite can be challenging to interpret because greigite can form at different periods significantly later than deposition, which then leads to anomalous paleomagnetic records and remagnetization [e.g., Jiang et al., 2001; Roberts and Weaver, 2005; Rowan and Roberts, 2006; Roberts et al., 2010; Sagnotti et al., 2010]. But greigite can also form during earliest burial and, thus, record an excellent paleomagnetic signal [e.g., Tric et al., 1991; Vasiliev et al., 2004, 2005; Hüsing et al., 2007] that matches well with the geomagnetic polarity time scale (GPTS) [Cande and Kent, 1995] and provides crucial age control for sedimentary sequences. Greigite is also useful for paleoenvironmental analysis. For example, greigite abundances from the Santa Barbara Basin have been demonstrated to reflect millennial-scale climate variability [Blanchet et al., 2009].

In sulfate-reducing sedimentary environments, greigite often forms as a precursor to pyrite (FeS2). As an intermediate phase that forms during pyritization, greigite is expected to fully convert to pyrite during early diagenetic sedimentary sulfate reduction [Berner, 1984]. Kao et al. [2004] demonstrated that when reactive iron is abundant and dissolved sulfide concentrations are low in sedimentary pore waters, pyritization can be arrested and greigite can be preserved, because dissolved sulfide (H2S, HS- ) is fully consumed by reactive
magnetostratigraphic dating and paleomagnetic reconstructions [Stoltz et al., 2007; Schumann et al., 2008; Chang et al., 2012; Larrasoana et al., 2012], and oceanic productivity [Roberts et al., 2011b; Yamazaki and Ikehara, 2012; Chang et al., 2013].

Unlike diagenetic greigite, which can remagnetize sediments or overprint primary remanences due to post-depositional growth, bacterial greigite should carry a syn-depositional paleomagnetic signal that is useful for magnetostratigraphic dating and paleomagnetic reconstructions [Vasiliev et al., 2008]. Compared to magnetite-producing MTB, greigite producers are less common in modern environments, although they have been identified in many sulfide-rich aquatic environments [e.g., Farina et al., 1990; Mann et al., 1990; Bazylinski et al., 1995; Pósfai et al., 1998a, 1998b; Simmons et al., 2004; Reitner et al., 2005; Wenter et al., 2009; Lefèvre et al., 2011; Wang et al., 2013]. Preservation of magnetite magnetofossils in sediments was expected to be common since the discovery of MTB in the 1970s [Kirschvink and Chang, 1984; Petersen et al., 1986; Stoltz et al., 1986; Hesse, 1994; Hounslow and Maher, 1996], and they are now being frequently reported [Kopp et al., 2007; Kopp and Kirschvink, 2008; Roberts et al., 2011b, 2012; Chang et al., 2012; Larrasoana et al., 2012; Yamazaki, 2012; Heslop et al., 2013]. In contrast, geological preservation of greigite magnetofossils is not well documented. Putative greigite magnetofossils have been reported from Miocene and Pliocene sediments from the Carpathian foredeep, Romania [Pósfai et al., 2001; Vasiliev et al., 2008]. Bacterial greigite has also been identified within Baltic Sea sapropels up to several thousand years in age [Reinholdsson et al., 2013]. Similar magnetic properties suggest that biogenic greigite may also magnetically dominate some Mediterranean sapropels [Roberts et al., 1999; Reinholdsson et al., 2013]. In order to assess the preservation potential of greigite magnetofossils in the geological record and to study greigite formation under changing environmental and climatic conditions, we use rock magnetic techniques, transmission electron microscope (TEM) observations, X-ray fluorescence (XRF), and cyclostratigraphy, to investigate Mio-Pliocene Black Sea sediments.

2. Geological Setting and Astronomical Tuning

Samples from several Mio-Pliocene sedimentary sections in the former Black Sea basin in Russia and Romania were selected for this study (Figure 1). Samples labeled “TK-XXX” and “TR-XXX” are from the Zheleznyi Rog (Iron Cape) section on the Black Sea margin of the Taman Peninsula, Russia. Standard paleomagnetic drill cores were taken at ~1–2 m stratigraphic intervals. This section is 500 m thick and covers ~5 Myr of Mio-Pliocene sedimentation in the Black Sea domain of the Paratethys Sea [Vasiliev et al., 2011]. We focus...
Figure 2. Chronology and climate stratigraphy for the Zheleznyi Rog section on the Black Sea coast, Taman Peninsula, Russia. (a) Field photograph with clear dark-light alternations related to climatic cyclicity. (b) Tentative astronomical tuning of the cyclic sediments to the La2004 astronomical solution [Laskar et al., 2004] and benthic δ¹⁸O records for the upper Miocene Ain el Beida section, northwestern Morocco [Van der Laan et al., 2005, 2006]. Numbers in Figure 2a correlate to the numbers in the lithostratigraphic log in Figure 2b. The green circle in (a) indicates people in the field as a scale.
on the upper 250 m of the section (Pontian and Kimmerian regional stages) [Krijgsman et al., 2010]. This is the time equivalent of the Messinian Salinity Crisis (MSC; 5.97–5.33 Ma) [Manzi et al., 2013; Roveri et al., 2014]. The base of the Pontian is characterized by a marine flooding event, which represents a reconnection of the Black Sea to the Mediterranean Sea [Krijgsman et al., 2010]. This event was followed by reestablishment of brackish conditions in the Black Sea basin and a major invasion of Pannonian species [Stoica et al., 2013; Grothe et al., 2014].

The Pontian sedimentary succession is characterized by distinct dark-light alternations (labeled A–D and 1–11 in Figure 2a), which indicate stable depositional conditions and a potential link to (astronomically) forced climate cyclicity. The alternations pass upward from dark bedded bituminous clay and marl, to diatomite couplets (A–C), to light and dark gray marls (D, 1–11). The diatomite couplets reflect increased marine conditions compared to the interbedded marls and represent diatom blooms at the transition between marine and freshwater conditions [Radionova and Golovina, 2011; Radionova et al., 2012]. Straightforward correlation to either obliquity or precession cannot be made; correlation to the astronomically tuned δ18O record of Van der Laan et al. [2005, 2006] provides a more straightforward pattern fit. This record from the Atlantic site of Morocco is the nearest oxygen isotope record not influenced by MSC evaporites and is taken as representative of regional climate. In our interpretation, light intervals correlate to cold/glacial stages and darker intervals to warmer/interglacial stages. The darker marls may represent a higher terrestrial component, related to higher river discharge during interglacial periods. In glacial periods, this is reduced and the “marine” calcite component is more dominant.

The geomagnetic reversal at 180 m is an age correlation tie-point that corresponds to the C3An.1n-C3r reversal at 6.033 Ma [Krijgsman et al., 2010; Vasiliev et al., 2011]. Our cyclostratigraphic correlation suggests a flooding age for the base of the Pontian to have been around 6.1 Ma, rather than the previously suggested 6.04 Ma [Krijgsman et al., 2010]. The majority of the Pontian was deposited during the long C3r reversed chron, which provides no additional magnetostratigraphic tie-points. The pattern of light beds 3–8 is, however, distinct and appears to correlate well with glacial δ18O stages TG 24 to TG 20 (Figure 2b). Overlying the Pontian marls is a distinct reddish interval that contains oolites/pisolites and nodular marls, which suggest high-energy coastal deposition. This reddish layer marks the base of the Kimmerian stage and is interpreted as the MSC climax related to glacial peaks TG 12 and TG 14, which recorded a sea level drop in the Black Sea basin [Krijgsman et al., 2010]. Faunal indicators suggest a change from brackish-marine conditions in the Pontian stages to a freshwater environment in the Kimmerian [Krijgsman et al., 2010], where no clear lithological cyclicity is observed.

Selected samples (“BD-XXX” and “RR-XXX”) from Pliocene Black Sea sediments of the Romanian Carpathians [Vasiliev et al., 2004, 2005] were also analyzed. These sediments, consisting of blue to gray sandstones, siltstones and clays, were deposited at high sedimentation rates (60–150 cm/kyr) in brackish environments. The samples are the same as those studied by Vasiliev et al. [2008]. Several synthetic greigite samples (“S-XXX”) [Chang et al., 2007, 2008] and magnetite magnetofossil-bearing sediment samples were analyzed to compare with results from greigite-bearing sediments. Sample “CD143-1-21” is a surface sediment sample (21 cm below the surface) from piston core CD143–55705 from the Oman margin [Rowan et al., 2009]. Sample “73C-11R1–40” is a pelagic carbonate from the Paleocene-Eocene thermal maximum from ODP Hole 738C, Southern Ocean [Larrasoana et al., 2012]. Sample “689D-11H6–21” is an early Oligocene pelagic carbonate from ODP Hole 689D, Southern Ocean [Roberts et al., 2012].

3. Methods

Room temperature hysteresis, isothermal remanent magnetization (IRM) acquisition, backfield demagnetization, and FORC measurements were performed using a Princeton Measurements Corporation MicroMag alternating gradient magnetometer (AGM) (Model 2900; noise level 2 × 10−9 Am−1) at the Paleomagnetic Laboratory of Utrecht University. Hysteresis loops were measured between ±1 T with a field step of 4 mT and a 150 ms averaging time. Hysteresis parameters, including the saturation magnetization (Ms), the saturation remanent magnetization (Mr), and coercivity (Br), were determined after paramagnetic slope correction. FORC diagrams [Pike et al., 1999; Roberts et al., 2000] were obtained with maximum applied fields of 1 T, field increments up to 0.4 mT, and averaging times of 100–300 ms. FORC diagrams were calculated using a program written by Tom Mullender at the Paleomagnetic Laboratory, Utrecht University. Data
representation close to the vertical axis of the FORC diagrams may be biased due to data extrapolation in this region. To avoid uncertain interpretation in this poorly defined part of the FORC space, it is not considered by the software. A smoothing factor (SF) between 3 and 5 was used. IRM acquisition curves were obtained by measuring 100–150 logarithmically spaced field steps up to a maximum field of 1 T on the AGM. IRM curves for some magnetically weak samples were measured using standard paleomagnetic core samples (~2 × 2.5 cm) on a robotized superconducting rock magnetometer (noise level 1–2 × 10⁻¹² Am²) at Utrecht University. Sixty data points were measured up to 700 mT. IRM acquisition curves were decomposed into coercivity components using the fitting protocol of Kruiver et al. (2001).

Thermomagnetic runs were measured in air with a modified horizontal translation-type Curie balance with a sensitivity of ~5 × 10⁻⁹ Am²² [Mullender et al, 1993] at Utrecht University. The applied field was cycled between 100 and 300 mT. Multiple heating and cooling cycles between room temperature, 200, 300, 350, 450, 620, and 700 °C were performed at a heating/cooling rate of 6 °C/min. Depending on sample magnetization, ~10–100 mg of sediment was used. A quartz glass holder and quartz wool were used to hold samples. Low-temperature magnetic properties were measured with a Quantum Design Magnetic Property Measurement System (MPMS; model XL7) at the Australian National University (ANU). For zero-field cooled (ZFC) and field-cooled (FC) curves, samples were cooled to 10 K in either zero-field or a 5 T field. At 10 K, a 5 T field was applied and then switched off to impart a low-temperature saturation IRM (SIRM), and the MPMS magnet was reset (the residual field after a magnet reset is ~200–300 μT). SIRM warming curves were measured during warming in zero-field. ZFC warming curves were acquired before the FC curves.

Ferromagnetic resonance (FMR) spectroscopy is used widely to detect biogenic magnetite chain signatures [e.g., Weiss et al, 2004; Kopp et al, 2006, 2007; Roberts et al, 2011b, 2012; Chang et al, 2014] and was employed to study our greigite-bearing samples. X-band FMR spectra were measured with a JEOL electron paramagnetic resonance (EPR) spectrometer at the Technical University of Munich or a Bruker Elexys EPR spectrometer at ANU. The 30–100 mg of sediment was measured at a frequency of 9.1–9.4 GHz and power of 0.6–2 mW. We used the same FMR parameters as those in other studies [Kopp et al, 2006; Roberts et al, 2011b, 2012].

Magnetic particles were extracted from bulk sediments by adapting the methods of Petersen et al [1986] and Vasiliev et al [2008]. Approximately 10 g of ground sediment was mixed with 300 mL of argon-purged demineralized water. Sodium polyphosphate (Na₄P₂O₇·10H₂O) was added to disperse the clays. The sediment solution was agitated using an ultrasonic bath for 30 min. The obtained sediment suspension was circled through a Frantz isodynamic magnetic separator for several hours. Magnetic extracts were then washed several times with demineralized water using a bar magnet. Magnetic extracts were viewed and analyzed using either a FEI Tecnai 20 FEG TEM at an acceleration voltage of 200 kV at Utrecht University, or a Philips CM300 TEM operated at 300 kV at ANU. Elemental compositions and crystal structures were analyzed using energy dispersive spectroscopy (EDS) and selected-area electron diffraction (SAED).

XRF analysis was carried out using a Thermo Scientific Niton XL3t XRF Analyzer at Utrecht University. Measurements were made on the top surface of standard paleomagnetic cylindrical samples. One flat surface of the cylinder was cleaned directly prior to data acquisition to ensure that data were obtained from fresh samples. Three measurements were averaged. Data that were significantly different from other measurements (e.g., when the X-ray beam interacted with large fossil shells) were excluded and measurements were repeated. Our XRF analysis is semiquantitative because no standard samples were used for calibration. Our purpose is to track elemental trends through the sequence, rather than absolute elemental concentrations. There are good correlations between large-scale elemental variations and magnetic properties, which are expected to reflect real sedimentary variations.

Samples from the Zheleznyi Rog section were taken at ~1–2 m stratigraphic intervals for paleomagnetic analysis [Krijgsman et al, 2010; Vasiliev et al, 2011]. Sister paleomagnetic samples were used for rock magnetic, XRF, and electron microscopic analyses. A total of ~150 samples from Taman (selected from the 0–230 m stratigraphic interval) were measured for hysteresis properties, IRM acquisition and backfield curves, and XRF elemental variations; ~40 were measured for FORC diagrams, ~20 were selected for thermomagnetic analyses, and 6 were selected for low-temperature magnetic analyses, FMR, and TEM observation. For samples from Romania, ~20 paleomagnetic drill core sister samples [Vasiliev et al, 2004, 2005, 2008] were selected for hysteresis, IRM, FORC, FMR, and thermomagnetic analyses.
4. Results

4.1. Rock Magnetism

4.1.1. Hysteresis Properties and FORC Diagrams

The studied samples have a wide range of hysteresis properties, mostly with single domain (SD) behavior (Figures 3a–3c). One type of hysteresis loop has relatively square shapes with $M_r/M_s$ ratios of $\sim 0.28-0.43$ and $B_r$ values of $\sim 10-25$ mT (Figure 3a). The other type has high $B_r$ values (mostly between 40 and 50 mT) and high $M_r/M_s$ values up to $\sim 0.7$, which are dominated by cubic magnetocrystalline anisotropy [Roberts, 1995] (Figure 3b). Some samples have intermediate $B_r$ and $M_r/M_s$ values mostly between these two types of hysteresis loops (Figure 3c). We classified the studied SD-dominated samples into three groups by their hysteresis and FORC characteristics: type-A (lower $B_r$ and square loops), type-B (higher $B_r$), and type-C (intermediate between type-A and type-B). We use this classification consistently throughout this paper. Hysteresis ratios for some samples are plotted after Day et al. [1977] in supporting information Figure S1.

High-resolution FORC measurements indicate variable coercivity and magnetostatic interaction distributions (Figures 3d–3f). Type-A samples (Figure 3d) have a dominant narrow ridge along the $B_r$ axis with negligible vertical spread. This FORC signature corresponds to that of noninteracting or weakly interacting SD particles [Pike et al., 1999; Egli et al., 2010]. Type-B samples have FORC diagrams (Figure 3e) with concentric contours with large vertical spread indicative of strong magnetostatic interactions [Pike et al., 1999; Roberts et al., 2000]. FORC diagrams for type-C samples have a mixture of type-A and type-B signatures: with both a central ridge and vertically spread concentric contours (Figure 3f). These samples, therefore, contain a variable mixture of noninteracting and interacting SD particles. FORC measurements for the same type-C samples measured with different numbers of FORCs but covering the same FORC region indicate that the central ridge signature is not clear in low-resolution FORC diagrams (Figures S2a and S2b), but it becomes more clearly evident in high-resolution FORC diagrams (Figures S2c and S2d).

4.1.2. IRM Acquisition and Demagnetization

Coercivity of remanence ($B_{c30}$) values, determined from backfield demagnetization measurements, range between $\sim 30$ and 70 mT. IRM acquisition curves for most samples can be fitted [Kruiver et al., 2001] with two main components (Figures 3g–3i). For type-A samples, one IRM component has a median field of $\sim 50-70$ mT and a small fitted dispersion parameter (DP) in the range of 0.20–0.22 (Figure 3g, magenta line). This component contributes $\sim 55$–80% of the total IRM. The other IRM component has a smaller median field ($\sim 25–36$ mT) and broader distribution (DP = $\sim 0.45–0.48$) (Figure 3g, green line). Type-B samples also have two major IRM components, one with a narrow distribution (Figure 3h, magenta line) and the other with a broad distribution (Figure 3h, green). The narrow IRM component for type-B samples contributes $\sim 60$–70% of the total IRM intensity. Compared to type-A samples, the narrow IRM component for type-B samples has higher median field (mostly between $\sim 75$ and 80 mT). DP values for this component are even smaller (mostly between 0.13 and 0.16). The other major component with broader distribution for type-B samples has median field values of $\sim 40–67$ mT, and DP values of $\sim 0.32–0.36$. Type-C samples (Figure 3i) have IRM components similar to type-B samples. All types of samples have an IRM component with DP < 0.22. The small low-field IRM component for all types of samples likely reflects deviations of the major IRM component from an ideal lognormal model distribution [Egli, 2004], to which we do not attribute physical meaning. This “skewed-to-the-left” behavior requires an extra component if fitting is restricted to symmetric distributions [Kruiver et al., 2001].

4.1.3. High and Low-Temperature Magnetic Properties

Thermomagnetic runs in air during different heating and cooling cycles indicate an irreversible decrease in magnetization between 200 and 400 °C with most pronounced drops in magnetization from $\sim 350$ to 420 °C in many samples (Figures 3j–3i). An irreversible decrease in magnetization between 200 and 420 °C is indicative of greigite [e.g., Reynolds et al., 1994; Roberts, 1995; Dekkers et al., 2000; Vasiliev et al., 2008; Chang et al., 2008; Roberts et al., 2011a] and is not characteristic of stoichiometric magnetite. For oxidized magnetite, such as magnetite with a maghemite (Fe₃O₄) coating, further oxidation occurs during heating in air, which causes irreversible magnetization loss. However, this loss is much less pronounced. Most samples also undergo a magnetization increase starting at $\sim 400$ °C, with a peak at $\sim 500$ °C, which indicates formation of a new magnetic mineral (most likely magnetite) due to oxidation of iron sulfides (usually pyrite) [Passier et al., 2001]. Variations in the magnitude of this high-temperature peak with respect to the magnetization...
loss below 400°C probably indicates variable concentrations of pyrite and greigite within the studied samples. Cooling from 700°C indicates that most of the newly formed magnetite transformed into hematite during heating.
We measured hysteresis properties and FORC diagrams for a range of samples at room temperature after heating to different temperatures and compare these with room temperature measurements without heating (Figure S3). After heating to 400°C, the characteristic noninteracting FORC signature for greigite-bearing sediments disappears (Figure S3a). Thermal decomposition of greigite below 400°C has been widely documented [e.g., Skinner et al., 1964; Reynolds et al., 1994; Roberts, 1995; Dekkers et al., 2000; Vasiliev et al., 2008; Chang et al., 2008; Roberts et al., 2011b]. In contrast, hysteresis loops and FORC diagrams for samples known to contain magnetite magnetofossils remain nearly the same after heating to 400°C (Figures S3b and S3c).

Low-temperature magnetic measurements indicate a nearly continuous decrease in remanence upon warming. No clear Verwey transition (T_V) is detected (Figures 3m–3o). This is consistent with the presence of greigite [Moskovitz et al., 1993; Roberts, 1995; Chang et al., 2009] and precludes the presence of significant magnetite within our samples. The large divergence between ZFC and FC curves below 35 K for samples from Taman (Figures 3m and 3o) is consistent with the magnetic behavior of siderite (FeCO_3) [Housen et al., 1996; Frederichs et al., 2003], which is a common authigenic mineral in reducing diagenetic sedimentary environments. This low-temperature discontinuity at 35 K is not due to the magnetic behavior of monoclinic pyrrhotite (Fe_S_3) [Dekkers et al., 1989; Rochette et al., 1990], as no discontinuity around 35 K is observed during low-temperature cycling of a room temperature SIRM.

### 4.2. FMR Spectroscopy

Room temperature FMR spectra for selected samples are presented in Figures 3p–3r. Most measured samples have a peak at a g value of 4.3 due to paramagnetic high-spin Fe^{3+}, and a characteristic Mn^{2+} sextet signal (labeled in Figures 3p–3r). The Mn^{2+} and paramagnetic Fe^{3+} signals are not of interest in this study and were therefore removed before calculating the FMR parameters. The residual FMR spectra for all types of samples have only one low-field peak and one broad high-field peak (Figures 3p–3r). These spectra contrast with those for intact biogenic magnetite [Weiss et al., 2004; Kopp et al., 2006, 2007; Roberts et al., 2011a, 2012; Larraoaria et al., 2012]. Particularly, they lack the multiple low-field absorption peaks and sharp high-field peak. These samples have g_{eff} of 2.0–2.3, \Delta B_{FWHM} values of 210–250 mT, asymmetric ratio A values close to 1, and z values larger than 0.3 (0.36–0.39). These FMR parameters do not fall within the regions for magnetite magnetosome chains [Weiss et al., 2004; Kopp et al., 2006].

### 4.3. TEM Observations

TEM observations were performed on magnetic extracts for both type-A and type-C samples (where a central ridge FORC feature is found) from the Black Sea sequence at Taman. Abundant iron sulfide nanocrystals are observed (Figures 4a–4c), which have been confirmed by EDS analysis (Figures 4j–4l). SAED patterns on selected extracted crystals are consistent with the crystal structure of greigite. Most crystals have octahedral, prismatic, round, or irregular shapes (Figure 4). Some greigite crystals have sizes and morphologies that are consistent with biogenic greigite crystals observed in living MTB, i.e., octahedral, cubo-octahedral, prismatic, and roundish, with a size range of ~20–100 nm and average size of ~60–75 nm [Farina et al., 1990; Mann et al., 1990; Bazylnski et al., 1995; Kasama et al., 2006; Muxworthy et al., 2013], which suggests a biogenic origin. Iron oxides (Figure 4d) are rare. This is consistent with bulk magnetic measurements, which indicate that greigite, rather than magnetite, is the main magnetic mineral in these sediments.

TEM-EDS analysis indicates the common presence of nickel (Figure 4f) within some larger iron sulfide crystals (100–500 nm). The ratio of Fe and Ni peak areas in TEM-EDS spectra is large (~38, 23, 11, and 5 for four crystals), which indicates variable Fe-Ni composition. In order to determine the Fe-Ni-bearing sulfide mineral phase, SAED patterns (Figures 4f–4l) were recorded on several representative Fe-Ni sulfide crystals, including the one in Figure 4e. The distinct crystal structure of greigite (cubic; space group Fd3m), pyrite (cubic; space group Pa3), and other common iron sulfide minerals (mackinawite: tetragonal P4/nmm; monoclinic pyrrhotite: A2/a) makes them readily distinguishable using SAED analysis. All measured diffraction patterns are consistent with those expected for the cubic crystal structure of greigite (Figures 4f–4i). Diagenetic iron-nickel sulfides have only been reported from Plio-Pleistocene marine marls from the Vrica section, Calabria, Italy [Van Velzen et al., 1993]. However, Van Velzen et al. [1993] could not provide crystal structure identification along with their microprobe analyses of this Fe-Ni-S phase, and they suggested monoclinic pyrrhotite as a possible mineral phase. Ni substitution in diagenetic magnetic minerals has not been confirmed by follow-up studies of the Vrica section [Roberts et al., 2010], and it has not been observed elsewhere in greigite [Roberts et al., 2011a]. Our TEM-based chemical and structural analysis confirms, for the first time, that...
diagenetic Fe-Ni sulfides have a ferrimagnetic greigite structure, although diagenetic formation of cation-doped greigite and its effects on the magnetization of greigite-bearing sediments need further investigation.

5. Discussion

5.1. Magnetic Identification of Biogenic Greigite and Paleomagnetic Implications

Biogenic magnetite produced by MTB has a range of distinct properties, such as characteristic crystal morphologies, narrow particle size distribution, chain structure, and SD magnetic behavior, which can be used to discriminate it from other types of magnetic minerals. Biogenic magnetite can be identified by direct TEM observations of magnetic mineral extracts and by a range of magnetic methods, including low-temperature magnetic tests [Moskowitz et al., 1993, 2008; Weiss et al., 2004; Chang et al., 2013], IRM analysis [Kruiver and Passier, 2001; Egli, 2004], FMR spectroscopy [Weiss et al., 2004; Kopp et al., 2006; Roberts et al., 2012], and FORC diagrams [Egli et al., 2010; Roberts et al., 2011b, 2012]. In contrast, detection of biogenic greigite is expected to be more difficult. First, biogenic greigite crystals tend to have less regular morphologies [e.g., Farina et al., 1990; Mann et al., 1990; Pósfai et al., 1998a, 1998b; Kasama et al., 2006] compared to
the distinct crystal morphologies of bacterial magnetite [Petersen et al., 1986; Stoltz et al., 1986; Roberts et al., 2011b; Chang et al., 2012; Larraoaia et al., 2012; Yamazaki, 2012]. Sedimentary diagenesis can also produce inorganic greigite crystals with a large grain-size spectrum [Pösfai et al., 2001; Rowan and Roberts, 2006; Roberts et al., 2011a] that includes the narrow size range of biogenic greigite crystals [Farina et al., 1990; Mann et al., 1990]. These factors make biogenic greigite crystal morphologies much less distinctive of biogenicity. Moreover, the often nonideal chain architecture of biogenic greigite [Farina et al., 1990; Mann et al., 1990; Pösfai et al., 1998a, 1998b; Kasama et al., 2006] compared to biogenic magnetite, and the absence of a low-temperature phase transition for greigite [Moskowitz et al., 1993; Roberts, 1995; Chang et al., 2009] mean that magnetic protocols for identifying biogenic magnetite may not be applicable to biogenic greigite. Additionally, extraction of magnetic minerals from lithified sedimentary rocks will cause significantly more disruption of magnetosome chain structures than in unconsolidated sediments and poorly cemented rocks.

Reinholdsson et al. [2013] observed that greigite magnetofossil-bearing Baltic sapropels have a FORC central ridge signature similar to that of biogenic magnetite. Biogenic magnetic minerals (both greigite and magnetite) preserved as isolated chains within sediment matrix are magnetically equivalent to noninteracting uniaxial SD particles. Such particle systems produce FORC distributions with a central ridge signature that has often been observed for intact biogenic magnetite chains [Egli et al., 2010; Roberts et al., 2011b, 2012]. In contrast, sedimentary diagenesis usually produces close-packed three-dimensional greigite clusters [e.g., Jiang et al., 2001; Roberts and Weaver, 2005; Rowan and Roberts, 2006; Sagnotti et al., 2010]. This distinct microstructure of diagenetic greigite results in strong magnetostatic interactions that produce FORC diagrams with significant vertical spread [Roberts et al., 2000, 2006; Rowan and Roberts, 2006; Chang et al., 2007; Vasiliev et al., 2007]. This type of FORC diagram has two other features: (1) the center of the distribution is shifted downward to negative $B_r$ values; and (2) there is a negative peak close to the $B_r$ axis in the lower quadrant. These signatures all indicate a strongly interacting SD magnetic particle assemblage [Pike et al., 1999; Roberts et al., 2000; Newell, 2005]. Such FORC signatures are observed for type-B and type-C samples that contain diagenetic greigite and are absent for type-A samples (biogenic origin). FORC diagrams, therefore, appear to be an excellent tool for discriminating biogenic from diagenetic greigite (Figures 3d–3f) [see also Reinholdsson et al., 2013]. We do not know the exact configuration of biogenic greigite crystals in sediments due to magnetic extraction-induced disruptions. Although fossilized biogenic greigite chains in sediments may not be well preserved, greigite magnetofossils cannot form close-packed particle clusters. Such distinct crystal arrangements, i.e., intact or disrupted biogenic greigite chains compared to diagenetic greigite in close-packed grain clusters, produce distinct magnetostatic interactions and FORC diagrams. FORC diagrams for some of our studied samples contain a central ridge (Figures 3d, 3f, and S1). This central ridge feature is unlikely to be due to initial growth of diagenetic greigite because early diagenetic greigite normally contains ultrafine particle clusters, with dominantly superparamagnetic properties and strong magnetostatic interactions [Rowan et al., 2009]. Such properties would not produce a central ridge feature. The central ridge signature also cannot be due to isolated greigite particles because greigite has cubic symmetry that would produce some vertical spread in FORC diagrams.

Our identification of biogenic greigite is supported by TEM observations that indicate the presence of abundant greigite nanocrystals, whose size and morphology are consistent with a biogenic origin (Figure 4). In addition, samples with a dominant central ridge feature (type-A) do not acquire a gyroremanent magnetization (GRM) during alternating field (AF) demagnetization [Krögsman et al., 2010; Vasiliev et al., 2011]. This GRM behavior contrasts with diagenetic greigite samples, which often contain closely packed greigite crystals that produce a bias field that enable GRM acquisition. No clear $T_s$ is detected (Figures 3m–3o), and thermal demagnetization of the natural remanent magnetization (NRM) on the same samples indicates that almost all remanence was removed below 420 °C [Krögsman et al., 2010; Vasiliev et al., 2011]. These observations indicate that there is no significant magnetite concentration in the studied samples.

FMR analysis provides another powerful tool for detecting biogenic magnetite chains [Weiss et al., 2004; Kopp et al., 2006]. We did not obtain a positive FMR test for our greigite-bearing samples (Figures 3p–3r). This may be due to the fact that greigite magnetosome chains often have less regularly arranged crystal orientations and chain alignment [Kasama et al., 2006]. Some greigite-producing MTB strains, such as magnetotactic rods and multicellular magnetotactic prokaryotes, have two or more adjacent chains. Significant departure from ideal linear chain structures—which is often found in intact greigite magnetosomes—will result in effective demagnetization and weaken the shape anisotropy.
would produce lower coercivities in biogenic greigite and cause a more gradual change in magnetization state, as has been confirmed by remanence measurements on single greigite-producing bacterial cells [Penningsa et al., 1995; Winkhofer et al., 2007] and hysteresis measurements on greigite magnetofossil-bearing sapropels [Reinholdsson et al., 2013].

Small DP values for IRM coercivity components have been used as an indicator of biogenic magnetite [Kruiver and Passier, 2001; Egli, 2004]. Biogenic greigite, which also has a narrow grain size distribution [Pösfaï et al., 2001; Kasama et al., 2006], is expected to produce small DP values, as observed in our type-A biogenic greigite samples (Figure 3g). In contrast to natural greigite samples (Figures 3g–3i), hydrothermally synthesized greigite samples [Chang et al., 2007, 2008] have one coercivity component with an ideal lognormal distribution and large DP values (0.31–0.33) (Figure S4). This is consistent with the fact that the hydrothermal synthesis method used produces particles with a wide range of grain sizes [Chang et al., 2007, 2008]. However, diagenetic SD greigite samples produce even smaller DP values (Figures 3h and 3i). The origin of such small DP values is not clear. It is possible that the nearly touching grains in diagenetic greigite clusters represent a strongly coupled magnetic system, where exchange coupling between neighboring crystals becomes important and results in collective switching of the magnetization, i.e., if one particle switches, all other particles switch due to exchange coupling. Nevertheless, our results indicate that a small DP value is not a unique indication of magnetosome crystals.

Despite the global occurrence of greigite-producing MTB in modern anoxic aquatic environments, fossilized greigite magnetosomes have not been widely reported in the geological record. In contrast, preservation of reactive iron and dissolved sulfate

5.2. Environmental and Climatic Control of Diagenetic Greigite Formation: Competition Between Reactive Iron and Dissolved Sulfate

Our integrated analysis, combining rock magnetism, XRF geochemistry, and sedimentary cyclicity, on Black Sea sediments from the Taman Peninsula enables unraveling of potential environmental control on diagenetic greigite formation (Figure 5). Diagenetic greigite usually forms as closely packed three-dimensional grain clusters in sediments, which produce a range of distinct rock magnetic properties that can be readily detected [Roberts et al., 2011a]. These magnetic characteristics include SD hysteresis properties, large coercivity values (often >35 mT), high Mr/Ms ratios (0.4), FORC diagrams indicative of strongly interacting assemblages, and GRM acquisition during AF demagnetization [Roberts et al., 2010]. Biogenic greigite (type-A) at Taman is less abundant than diagenetic greigite, and is mostly identified within the 100–120 m interval (Figure 5). Diagenetic greigite (both type-B and type-C) occurs throughout the section, and
is common in clay-rich dark layers and absent in carbonate-rich light layers (Figures 2 and 5). We use $B_c > 35$ mT and $M_t$ peaks as indicators of significant diagenetic greigite concentrations (horizontal bars; Figure 5). Biogenic greigite occurrences do not affect our paleoenvironmental interpretation for diagenetic greigite formation because diagenetic greigite typically has higher $B_c$ and $M_t$ values than biogenic greigite (Figures 3a–3f) [Reinholdsson et al., 2013]. Within zones 2 and 3, good correlations between diagenetic greigite concentration, lithology, and elemental abundances are identified (horizontal bars). In zone 3, diagenetic greigite concentration correlates to low Rb/Sr and high Ca/Al ratios (gray bars). In zone 2, diagenetic greigite spikes occur at intervals with high Rb/Sr and low Ca/Al (blue bars). Between 150 and 230 m (zone 1), samples are magnetically weak and lack any indication of greigite. The climatic index TG is from Van der Laan et al. [2005, 2006], which is the same as in Figure 2.

Stratigraphic elemental variations, expressed as relative concentrations, indicate two distinct zones (zone 3 versus zones 1 and 2 in Figure 5) separated by the reddish layer associated with the MSC climax at ~89–94 m (Figures 2 and S6). Across the red layer, a distinct switch in some elemental contents and ratios (Ca, Ti, Ca/Al, and Rb/Sr; Figures 5 and S6) indicates a significant environmental change. In zone 1 (~150–230 m), samples are magnetically weak (dominantly paramagnetic) and lack diagenetic greigite. Zone 2 (below the reddish layer) is characterized by generally high Ca/Al ratios and low Rb/Sr, compared to zone 3. Contrasting diagenetic greigite concentrations and geochemical parameters occur in zones 2 and 3. In zone 2, blue bars in Figure 5 indicate high $B_c$ and $M_t$ values (which indicate significant diagenetic greigite concentrations) correlate with low Ca/Al (i.e., weaker biogenic activity) and S/Al (i.e., less sulfidic sulfur) ratios, but high Rb/Sr (stronger continental weathering and more terrigenous input) [Blanchet et al., 2009]. In zone 3, high $B_c$ and $M_t$ values are indicated with gray shading in Figure 5, where they correlate with high Ca/Al and S/Al ratios, but low Rb/Sr.
In sulfate-reducing sedimentary environments, greigite forms as a precursor to pyrite through reaction of detrital iron-bearing minerals with $\text{H}_2\text{S}$ (produced by sulfate-reducing bacteria). These reactions are often limited by three principal factors: (1) availability of dissolved sulfate in sedimentary pore waters, (2) availability and reactivity of organic matter for microbial metabolism, and (3) reactive iron released from detrital minerals [Berner, 1984; Roberts et al., 1996; Roberts and Weaver, 2005]. Kao et al. [2004] demonstrated that $\text{HS}^-$ released by microbes can be effectively removed by high dissolved iron activity (i.e., from abundant reactive detrital iron-bearing minerals or coatings from which $\text{Fe}^{2+}$ dissolves), so that the pathway to pyrite formation is arrested and intermediate greigite is preserved. A similar mechanism was invoked by Blanchet et al. [2009] to explain the formation and preservation of diagenetic greigite in rapidly deposited marine sediments from the Santa Barbara Basin, where diagenetic greigite occurs in terrigenous-rich and organic-poor sedimentary layers. In such a scenario, diagenetic greigite peaks should correlate with high $\text{Rb}/\text{Sr}$ and low $\text{Ca}/\text{Al}$ ratios (stronger continental detrital contribution and weaker biogenic activity). This pattern is observed in zone 2 of the studied sediments (Figure 5, blue bars), where diagenetic greigite formation appears to have been controlled mainly by terrigenous input. In marine sedimentary environments, sulfate is abundant and is readily used by microbes to produce $\text{H}_2\text{S}$ (or $\text{HS}^-$). However, diagenetic greigite peaks also occur in intervals with lower continental detrital inputs in zone 3 (i.e., low $\text{Rb}/\text{Sr}$ and high $\text{Ca}/\text{Al}$). In lake environments, detrital iron-bearing minerals and reactive organic matter are often abundant and, therefore, should not limit diagenetic greigite formation, but dissolved sulfate concentrations are typically low [Roberts et al., 1996]. It has been suggested that the key factor for diagenetic greigite formation in lacustrine environments is dissolved sulfate availability, with greigite preservation being favored by sulfate consumption before reaction to pyrite is completed [Roberts et al., 1996]. This appears to have been the case in zone 3, which is supported by thermomagnetic data (Figure S7), where samples from the weakly magnetized interval (i.e., low $M_s$ and $B_c$ values) in zone 3 contain a dominant paramagnetic signal with no indication of pyrite (i.e., a pyrite peak at $\sim 500^\circ\text{C}$ is absent [Passier et al., 2001]; Figure S7). In contrast, weak samples in zones 1 and 2 often have a pronounced pyrite peak at $\sim 500^\circ\text{C}$ (Figure S7). This is also consistent with the observation that diagenetic greigite peaks correlate with $\text{S}/\text{Al}$ peaks.

Paleontological analysis of endemic Paratethys faunal assemblages (molluscs and ostracods) from the studied Taman section indicates an evolution from a marine to a lake environment across the MSC climax [Krijgsman et al., 2010]. This paleoenvironmental evolution is consistent with our interpretation of conditions that favored diagenetic greigite formation. Prior to the MSC climax, significant greigite formation probably coincided with climatic changes or sea level drops during the MSC. During this interval, diagenetic greigite formation occurred in warm/wet climatic periods (Figures 2 and 5) and was probably prompted by terrigenous enrichments in reactive iron supply driven by enhanced continental weathering around the Black Sea. After the MSC climax, the Black Sea basin was isolated and disconnected from the Mediterranean and became a large lake [Krijgsman et al., 2010]. During this period, diagenetic greigite enhancement was due to limnological conditions that caused small amounts of dissolved sulfide to react with abundant iron to cause greigite preservation. In summary, we observe contrasting environmental controls on diagenetic greigite formation between marine and lacustrine environments in the same sedimentary sequence. This supports a previous suggestion that diagenetic iron sulfides, including greigite, could be useful paleosalinity indicators [Berner et al., 1979].

6. Conclusions

We provide evidence for biogenic greigite preservation from Black Sea sediments in Russia and Romania. Biogenic greigite has magnetic properties that enable discrimination from other types of magnetic minerals, particularly diagenetic greigite. FORC analysis is the most useful method for identifying biogenic greigite: the central ridge signature of biogenic greigite contrasts with vertically distributed signatures for diagenetic greigite. High-temperature magnetic properties enable discrimination of greigite from magnetite due to thermal decomposition of greigite below 420 $^\circ\text{C}$. Our analysis suggests that preservation of greigite magnetofossils is potentially widespread in ancient sediments. Their high preservation potential is important for understanding the paleomagnetic reliability of greigite-bearing sediments, which can be complicated by remagnetizations. In addition, our TEM analysis enabled identification of the common presence of nickel-doped iron sulfides, which we confirm to have a greigite structure.
Integrated rock magnetic, geochemical and cyclostratigraphic analysis of a sedimentary sequence from the Black Sea basin provides an improved age model for understanding the evolution of the Black Sea, particularly during the Messinian Salinity Crisis. We observe correlation between diagenetic greigite concentrations and certain geochemical features, which demonstrate an environmental control on diagenetic greigite formation. Prior to the MSC climax, the Black Sea basin in Taman was dominantly a marine environment that became progressively disconnected from the Mediterranean Sea. Greigite formation within this interval appeared to have been climatologically controlled with greigite layers occurring during warm/wet periods, which are interpreted to have been associated with enhanced terrigenous input. After the MSC climax, the Black Sea basin became isolated from the Mediterranean Sea and a freshwater environment dominated. This environmental change altered the relative abundance of detrital iron minerals, organic matter and dissolved pore water sulfate that resulted in different authigenic greigite formation patterns. Our studies demonstrate the usefulness of magnetic analysis, particularly greigite detection, for paleoenvironmental reconstructions.

**References**


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