Environmental control on the occurrence of high-coercivity magnetic minerals and formation of iron sulfides in a 640 ka sediment sequence from Lake Ohrid (Balkans)

Janna Just1,2, Norbert R. Nowaczyk3, Leonardo Sagnotti4, Alexander Francke1, Hendrik Vogel5, Jack H. Lacey6, and Bernd Wagner1,2

1Institute of Geology and Mineralogy, University of Cologne, Cologne, Germany
2Collaborative Research Centre 806 – Our Way to Europe, University of Cologne, Cologne, Germany
3Helmholtz Centre Potsdam, GFZ German Research Centre for Geosciences, Potsdam, Germany
4Istituto Nazionale di Geofisica e Vulcanologia, Rome, Italy
5Institute of Geological Sciences & Oeschger Centre for Climate Change Research, University of Bern, Bern, Switzerland
6NERC Isotope Geosciences Facilities, British Geological Survey, Nottingham, UK

Correspondence to: Janna Just (janna.just@uni-koeln.de)

Received: 31 July 2015 – Published in Biogeosciences Discuss.: 28 August 2015
Revised: 25 January 2016 – Accepted: 8 February 2016 – Published: 8 April 2016

Abstract. The bulk magnetic mineral record from Lake Ohrid, spanning the past 637 kyr, reflects large-scale shifts in hydrological conditions, and, superimposed, a strong signal of environmental conditions on glacial–interglacial and millennial timescales. A shift in the formation of early diagenetic ferrimagnetic iron sulfides to siderites is observed around 320 ka. This change is probably associated with variable availability of sulfide in the pore water. We propose that sulfate concentrations were significantly higher before ∼320 ka, due to either a higher sulfate flux or lower dilution of lake sulfate due to a smaller water volume. Diagenetic iron minerals appear more abundant during glacials, which are generally characterized by higher Fe/Ca ratios in the sediments.

While in the lower part of the core the ferrimagnetic sulfide signal overprints the primary detrital magnetic signal, the upper part of the core is dominated by variable proportions of high- to low-coercivity iron oxides. Glacial sediments are characterized by high concentration of high-coercivity magnetic minerals (hematite, goethite), which relate to enhanced erosion of soils that had formed during preceding interglacials. Superimposed on the glacial–interglacial behavior are millennial-scale oscillations in the magnetic mineral composition that parallel variations in summer insolation. Like the processes on glacial–interglacial timescales, low summer insolation and a retreat in vegetation resulted in enhanced erosion of soil material. Our study highlights that rock-magnetic studies, in concert with geochemical and sedimentological investigations, provide a multi-level contribution to environmental reconstructions, since the magnetic properties can mirror both environmental conditions on land and intra-lake processes.

1 Introduction

Rock-magnetic properties of sediments can be used to reveal changing input of the lithogenic fraction and can therefore serve as records of past environmental change. Variations in the concentration of magnetic minerals provide information on changing export of terrigenous sediments from land (DeMenocal et al., 1991; Just et al., 2012; Maher, 2011; Reynolds and King, 1995). Furthermore, variations in the composition of the magnetic mineral assemblages can be used for detecting changes in terrestrial climatic conditions, e.g., weathering and soil formation (Hu et al., 2015; Kämpf and Schwertmann, 1983; Larrasoña et al., 2015; Lyons et
al., 2010; Maher and Thompson, 1992). However, in addition to the detrital magnetic inventory, magnetic minerals form in the course of syn- and post-sedimentary processes. Magnetotactic bacteria living in the oxic–anoxic transition zone in the topmost sediments utilize magnetic minerals, and can produce either magnetite or greigite intracellularly (Egli, 2004b; Roberts et al., 2012; Snowball et al., 2002; Vali et al., 1989). Furthermore, iron-reducing bacteria may induce the authigenic precipitation of magnetite (e.g., Bell et al., 1987; Frankel and Bazylinski, 2003). On the other hand, the primary magnetic mineral assemblage of detrital origin is often overprinted by post-depositional alteration (Hounslow and Maher, 1999; Nowaczyk et al., 2013; Roberts et al., 1996). The latter results from changing redox conditions at the lake/sea floor and in subsurface sediments leading to dissolution of iron oxides and formation of ferrimagnetic iron sulfides, such as greigite and pyrrhotite (Demory et al., 2005; Froelich et al., 1979; Karlin and Levi, 1983; Rowan et al., 2009; Sagnotti, 2007; Vasiliev et al., 2007) or paramagnetic minerals, such as pyrite, siderite, and vivianite (Dong et al., 2000; Karlin and Levi, 1983).

In addition to the vast number of studies on magnetic minerals in marine sediments, lake sediments can provide valuable information on terrestrial and lacustrine environmental conditions (e.g., Frank et al., 2002; Nowaczyk et al., 2002; Peck et al., 1994, 2004; Roberts et al., 1996; Snowball, 1993; Snowball et al., 1999; Wang et al., 2008). Depending on the trophic state of the lake, water depth, and stratification, oxygen supply is often limited and may lead to excessive dissolution of iron oxides (e.g., Demory et al., 2005; Frank et al., 2013; Nowaczyk et al., 2013; Snowball et al., 1999). In the course of early diagenesis, sulfate is reduced during the process of organic matter degradation in the sulfidic zone (Froelich et al., 1979). H$_2$S species can react with accessible iron and form iron sulfides. Among them, pyrrhotite and greigite are particularly important for rock and paleomagnetic studies (Roberts et al., 2011a; Sagnotti, 2007). As these secondary magnetic iron minerals acquire a remanent magnetization iron sulfides can bias the primary magnetic signals (Ron et al., 2007), in general carried by detrital (titanomagnetite. Although pyrrhotite and greigite may form as early diagenetic metastable phases during the chemical reaction pathway to pyrite, studies have shown that they may be preserved if the concentration of organic matter, and consequently organic-bound sulfur, is low and pyritization is hampered by Fe excess (Blanchet et al., 2009; Kao et al., 2004; Roberts et al., 1996; Skinner et al., 1964). Cyclic preservation of greigite in a Pliocene lacustrine sequence from Lake Qinghai (China) was recently linked to periods of high lake levels and humid climate that resulted in oxygen depletion of the bottom water (Fu et al., 2015).

Compared to the ocean, lakes generally contain lower sulfate concentrations, and therefore sulfate may be exhausted at shallow depths in the sediment or even in the water column. Below the sulfidic zone, methanogenesis is the most important process for the degradation of reactive organic matter (Capone and Kiene, 1988; Roberts, 2015). Here, Fe species may react to form iron phosphate (vivianite) or iron carbonate ( siderite; Berner, 1981; Roberts, 2015). In addition to the “early” diagenetic processes, concurring with organic matter degradation, authigenic Fe minerals can form at a late diagenetic phase. Such precipitation and dissolution processes occur mainly as a result of a variety of different mechanisms associated with the migration of mineralized fluids and with changes in the pore water chemistry, which disrupt the steady-state diagenetic progression (Roberts and Weaver, 2005; Sagnotti et al., 2005).

Because of the imprint of these interacting processes, rock-magnetic properties of lake sediments can provide a suite of environmental information. These range from variations in the external supply of magnetic detrital mineral phases to changes in lake water oxygenation and sulfate supply and in conditions favorable for magnetotactic bacteria. The sediments from Lake Ohrid on the Balkan Peninsula (Fig. 1) provide an excellent opportunity to study these processes over several glacial–interglacial cycles. Lake Ohrid is the oldest lake in Europe, dating back to >1.2 Ma (Wagner et al., 2014). Its sediments consist of lacustrine carbonates (mostly calcite), biogenic silica, and lithogenic compounds (Francke et al., 2015). Lake Ohrid is an oligotrophic lake; a complete overturn and deep mixing occur only every few years, while the upper 200 m of the water column is mixed every year during winter (e.g., Matzinger et al., 2007). Temperature variations probably had a strong influence on shallow and deep mixing of Lake Ohrid during past glacial and interglacial cycles (Vogel et al., 2010a; Wagner et al., 2009); however, to date there is no quantitative estimate of this effect.

Terrestrial element concentrations and gamma ray intensities of Lake Ohrid sediments mirror phases of enhanced erosion in the catchment on glacial–interglacial timescales (Baumgarten et al., 2015; Francke et al., 2015). Geochemical variations in a core from northeastern Lake Ohrid (Fig. 1) are indicative of changes in sediment dynamics over the last ~140 kyr. Here, the increasing deposition of Cr-rich minerals, most likely originating from the western flanks of Lake Ohrid, are thought to result from either enhanced erosion of soils or stronger wind activity, inducing changes in surface water circulation (Vogel et al., 2010a).

This first study on the magnetic properties of the Lake Ohrid “DEEP” Site (ICDP 5045-1) focuses on observations of changing magnetic mineralogy and possible implications for lacustrine and terrestrial environmental conditions. We aim to identify primary detrital magnetic minerals, and evaluate whether these reveal changing environmental conditions in the catchment, beyond the observed general pattern of higher (lower) terrigenous input during glacial (interglacial). Secondly, we discuss the occurrence of (early) diagenetic minerals to develop a working hypothesis concerning changes in redox conditions and shifts in the chemical and hydrological environment in the lake. To address these
objectives, we jointly investigated mineral magnetic properties and organic proxies, X-ray fluorescence (XRF) data (Francke et al., 2015) and Fourier transform infrared spectroscopy (FTIRS; Lacey et al., 2016).

2 Study area and materials

Lake Ohrid (45°54'N, 38°20'E; Fig. 1) is located at the boundary between Albania and the Former Yugoslav Republic of Macedonia at an altitude of 690 m above sea level. It is ∼30 km long and ∼15 km wide, and has a maximum water depth of 293 m. It is flanked by high mountain ranges in the west (Jurassic ophiolites and Triassic carbonates) and east (Triassic carbonates), and an alluvial plain in the north (Fig. 1, Hoffmann et al., 2010, and references therein; Vogel et al., 2010b). Modern vegetation is dominated by deciduous forest (Lézine et al., 2010).

At present, there are two major rivers draining into Lake Ohrid, the Sateska in the north and Cerava in the south, bringing detrital sediments to the lake. Freshwater is supplied to the lake by rivers (25%), direct precipitation (25%) and karst aquifers (50%). The karst aquifers are fed by mountain range precipitation and from the neighboring higher-altitude (849 m above sea level) Lake Prespa. The gain in freshwater is balanced by the drainage through the river Crn Drim (accounting for 60% of water loss) and evaporation (∼40%) of lake water (Matzinger et al., 2006a, b; Wagner et al., 2014).

Maximum precipitation occurs during winter, and air temperatures at present range between −5 and 31 °C (Popovska and Bonacci, 2007). Ohrid is an oligotrophic lake with maximum productivity during summer (Matzinger et al., 2007; Wagner et al., 2014). However, at present, it shows indications for the onset of eutrophication (e.g., Matzinger et al., 2007).

Six holes were drilled at the “DEEP” site (ICDP Site 5045-1) as part of the SCOPSCO project (Scientific Collaboration on Past Speciation Conditions in Lake Ohrid) in summer 2013 down to a maximum sediment penetration of 569 m (Fig. 1b). The recovered sedimentary sequence is thought to cover more than 1.2 Myr (Wagner et al., 2014). A composite profile was constructed and an age model was developed by Francke et al. (2015) which is based on 11 teprostratigraphic tie points (Leicher et al., 2015) and correlating geochemical proxies to orbital parameters (using an age uncertainty of ±2000 years). Here, we analyzed the upper 247 m of the composite profile. The age model reveals that the analyzed interval spans the past 637 kyr.

3 Methods

3.1 Rock-magnetic measurements

We sampled the composite profile at 50 cm (0–100 m) and 48 cm (100–247 m) intervals – in total 500 samples – using 6.2 cm$^3$ oriented plastic boxes. Magnetic susceptibility $\chi$ was measured using an AGICO MFK-1 susceptometer. Natural and artificial remanence parameters were measured using a 2G Enterprises 755 superconducting long-core rock magnetometer with an in-line tri-axial alternating field (AF) demagnetizer. The natural remanent magnetization (NRM) was demagnetized in 10 incremental steps of up to 100 mT AF peak amplitude. Anhysteretic remanent magnetization (ARM), a proxy for fine-grained, mostly single domain (SD) and pseudosingle domain (PSD) magnetite (King et al., 1982), was imparted with a single-axis 2G 600 AF demagnetizer by using 100 mT AF and 50 μT DC field. Subsequently, ARM was AF-demagnetized at fields of 10, 20, 30, 40, 50, and 65 mT. The median destructive field
(MDF\textsubscript{ARM}), defined as the field required to decrease ARM by 50\%, was calculated. This parameter is indicative of the coercivity within the fine ferrimagnetic mineral fraction.

Isothermal remanent magnetization (IRM), which depends on the magnetic mineral mixture in the samples, was induced using a 2G 660 pulse magnetizer applying a 1500 mT DC peak field and a 200 mT reversed field. The ratio of $x_{\text{ARM}}$ to saturation IRM ($x_{\text{ARM}} / \text{SIRM}$) serves as a proxy for the magnetic grain size. Furthermore, hard IRM (HIRM), reflecting the contribution of high-coercivity magnetic minerals to SIRM, was calculated using the equation

$$\text{HIRM} = 0.5(\text{SIRM} + \text{IRM}(-200\text{mT})).$$

Additionally, the $S$ ratio, calculated as

$$S = 0.5 \times (1 - (\text{IRM} - 200\text{mT} / \text{SIRM})),$$

serves as a proxy for the proportion of high- (e.g., hematite + goethite, $0 < S < 1$) to low-coercivity (magnetite, greigite) magnetic minerals ($0 < S \leq 1$).

Moreover, we calculated SIRM / $x$, which is often observed to be elevated in the presence of greigite (e.g., Maher and Thompson, 1999; Nowaczyk et al., 2012; Snowball and Thompson, 1990). Another characteristic behavior of greigite is that it acquires a so-called gyro-remanent magnetization (GRM). To further quantify the possible imprint of greigite, we calculated the ratio between the differences of final remanent magnetization (FRM) at 100 mT AF peak amplitude and minimum magnetization (MRM) during NRM demagnetization, and the difference of NRM and MRM according to (Fu et al., 2008)

$$\Delta\text{GRM} / \Delta\text{NRM} = (\text{FRM-MRM}) / (\text{NRM-MRM}).$$

To account for down-core increasing sediment compaction, magnetic concentration parameters were mass-normalized using the dry bulk density. The latter data were available at 4 cm intervals and interpolated to the depths of magnetic samples.

### 3.2 Cluster analysis

We performed fuzzy c-means cluster analysis for a suite of data that are indicative of the magnetic mineralogy and magnetic granulometry. To achieve symmetric distributions of the suite of data, we performed data ln (natural logarithm) transformations, except for $\Delta\text{GRM} / \Delta\text{NRM}$ and the $S$ ratio. The latter parameters show a J-shaped distribution that cannot be transformed into a normal distribution. All data sets were standardized before clustering.

### 3.3 Scanning electron microscopy of magnetic extracts

Six samples that are characterized by divergent magnetic properties were selected for scanning electron microscopy (SEM) analyses. Magnetic extracts were prepared following the protocol of Nowaczyk (2011). A total of 2 cm$^3$ of sediment was dispersed in 60 mL of alcohol and Na solution was added. The solutions was put in an ultrasonic bath for 10 min. A rare-earth magnet inside a plastic hose was submerged into the solution. Minerals attracted to the hose were washed into a fresh vial. The procedure was repeated twice to obtain a clean extract. Particles in the alcohol solution were then concentrated and a drop of the solute was placed on an SEM stub and sputtered with carbon.

Magnetic extracts were analyzed using a Carl Zeiss SMT Ultra 55 Plus SEM. Images were obtained using the secondary and back scatter electron beams of single particles. To obtain the chemical composition of the analyzed particles, energy dispersive spectroscopy was performed at energy levels of 20 keV.

### 3.4 Geochemical and mineralogical data

Total organic carbon (TOC), total inorganic carbon (TIC) and X-ray fluorescence (XRF) data, measured by Francke et al. (2015), are used for discussion of rock-magnetic data. XRF scanning was carried out at 2.5 mm resolution and with an integration time of 10 s using an ITRAX core scanner equipped with a chromium X-ray source. Total carbon (TC) and total TIC were analyzed at 16 cm resolution. For TIC, homogenized and dispersed sediments were treated with phosphoric acid. TC and TIC was measured in the form of released CO$_2$ after combustion at 900 and 160°C, respectively, using a DIMATOC 100. Total organic carbon (TOC) content was calculated from the mass difference between TC and TIC. In addition, we show relative changes in siderite (FeCO$_3$) concentration determined from infrared (IR) spectra. IR spectra were measured on a Bruker Vertex 70 Fourier transform infrared (FTIR) spectrometer at the Institute of Geological Sciences at the University of Bern. Details on measurement set up and data evaluation procedures are outlined in Lacey et al. (2016).

### 4 Results

#### 4.1 Rock magnetism

In the magnetic properties, a major change is visible around ca. 320 ka corresponding to the transition between Marine Isotope Stage (MIS) 9 and MIS 8. Below this transition the $S$ ratio is rather constant but SIRM shows large amplitudinal change, while above the transition SIRM is relatively constant but compositional magnetic proxies, e.g. the $S$ ratio, show interglacial–glacial variability. We therefore divided the record into two magnetic units (Fig. 2).

#### 4.2 MIS 16–MIS 9 (unit 2)

Except for a stable $S$ ratio (Fig. 2e), this unit is characterized by high-amplitude variations on glacial–interglacial
timescales. Susceptibility (Fig. 2b) and SIRM (Fig. 2c) and HIRM (Fig. 2d) are elevated during glacial, while $\chi_{\text{ARM}} / \chi$ (Fig. 2g) and $\chi_{\text{ARM}} / \chi$ (Fig. 2h) decrease and the MDF$_{\text{ARM}}$ (Fig. 2f) rises to fields of up to 50 mT. Increasing $\Delta\text{GRM} / \Delta\text{NRM}$ (Fig. 2j) is often associated with interglacial–glacial transitions, as well as early glacial. Moreover, short-lasting spikes of GRM acquisition during interglacials are accompanied by brief increases in Fe/Ca ratios (Fig. 2k, e.g., during MIS 15, 13, 11). In contrast, SIRM / $\chi$ (Fig. 2i) shows maximum values at the end of the glacial (MIS10, MIS12), when GRM acquisition is mostly low.

4.3 MIS 8–MIS 1 (unit 1)

Compared to unit 2, glacial–interglacial variations are expressed through different proxies. SIRM / $\chi$ (Fig. 2i) is low and $\Delta\text{GRM} / \Delta\text{NRM}$ (Fig. 2j) is mostly zero. SIRM (Fig. 2c) and the MDF$_{\text{ARM}}$ (Fig. 2f) are relatively stable, while sus-
ceptibility (Fig. 2b) and HIRM (Fig. 2d) show glacial–interglacial cyclicity. \(\chi_{\text{ARM}} / \chi\) (Fig. 2e) is lower during glacials, in concert with higher TOC (Fig. 2f) and lower Fe / Ca ratios (Fig. 2k). An exception to this glacial–interglacial behavior is observed in the uppermost part of unit 1, where TOC increases and Fe / Ca decreases. In contrast to the relationship observed for the rest of the unit, the intensity of the magnetic concentration parameters (\(\chi / \chi_{\text{SIRM}}\), Fig. 2b, c) increases, and the \(S\) ratio rises to high values. Also, the MDF_{\text{ARM}} and \(\chi_{\text{ARM}} / \chi\) and \(\chi_{\text{ARM}} / \chi_{\text{SIRM}}\) rise to maximum values.

4.4 SEM observations

We analyzed samples that are characterized by diverging magnetic properties (see Table 1). Detrital titanomagnetites and chromium iron oxides are present in almost all analyzed samples (Fig. 3a, b, d). Magnetic extracts from the uppermost part of the unit contain relatively high proportions of siderite (Fig. 3g), whereas iron sulfides are abundant in samples from the lower part (Fig. 3c–f, h).

All analyzed samples in unit 2 are characterized by a high MDF_{\text{ARM}} (> 45 mT). Of those, two samples acquired a significant GRM (> 68 %), but had relatively low SIRM / \(\chi\), while the other two samples have the most extreme SIRM / \(\chi\) and insignificant GRM acquisition (Table 1). The high GRM samples contain mixtures of mixtures of titanomagnetites and fine-grained and microcrystalline iron sulfides, which most likely correspond to greigite (Fig. 3c, d). Also, the low GRM samples contained fine-grained iron sulfides and idiomorphic greigite crystals but, additionally, large iron sulfide nodules (Fig. 3e, f, h) that apparently have a higher Fe : S ratio compared to the finer-grained greigite (data not shown). Although the number of investigated SEM samples is relatively low, the results suggest that a high MDF_{\text{ARM}} appears closely related to or at least accompanied by the general presence of iron sulfides, while maximum SIRM / \(\chi\) is associated with the coarse-grained sulfide nodules. GRM is quite strong when microcrystalline or individual greigite crystals occur.

5 Discussion

5.1 Magnetic proxy evaluation

Many samples from unit 2 have high values in MDF_{\text{ARM}}, SIRM / \(\chi\) and \(\Delta\chi_{\text{ARM}} / \Delta\chi_{\text{NRM}}\) (Fig. 2f, i, j), which are often associated with the presence of greigite (e.g., Fu et al., 2008; Peters and Dekkers, 2003; Rowan et al., 2009). It is important to note that these parameters can sometimes provide ambiguous results; high SIRM / \(\chi\) is not always accompanied by high \(\Delta\chi_{\text{ARM}} / \Delta\chi_{\text{NRM}}\); however, MDFs are high if any of the former proxies are elevated. Missing GRM acquisition of greigite bearing samples has been reported in other studies (Roberts et al., 1998; Sagnotti et al., 2005), and could be related to grain-size effects, because only fine-grained (SD) greigite acquires GRM. As observed in the images of the magnetic extracts, coarse-grained iron sulfides are abundant in samples with low GRM and high SIRM / \(\chi\) and we therefore hypothesize that greigite either grew to too coarse grain size for GRM acquisition or was transformed into other iron sulfides, e.g., pyrrhotite, which is also characterized by high SIRM / \(\chi\) (Maher and Thompson, 1999). Variations on interglacial–glacial timescales within the upper unit are the second significant feature in the rock-magnetic record. The variations could stem from compositional changes in the magnetic mineral supply to the lake, but they could also result from selective dissolution of low-coercivity magnetic minerals. To further evaluate the rock-magnetic properties, we show cross plots of selected parameters (Fig. 4).

The \(\chi_{\text{ARM}} / \chi_{\text{SIRM}}\) ratio (Fig. 2g) is often utilized as a proxy for magnetic grain size, but it can be biased if SIRM is dominated by high-coercivity magnetic minerals (HIRM). However, HIRM is more than 2 orders of magnitude smaller than SIRM (Fig. 2), and thus SIRM can be applied as a proxy for the concentration of the low-coercivity fractions (e.g., magnetite and greigite).

For non-GRM samples, low \(S\) ratios are associated with low \(\chi_{\text{ARM}} / \chi_{\text{SIRM}}\) (Fig. 4a), implying that a shift towards high-coercivity minerals is accompanied by a coarsening of the low-coercivity fraction. In contrast, GRM samples have high \(S\) ratios but low \(\chi_{\text{ARM}} / \chi_{\text{SIRM}}\) values, indicating that in combination these parameters are valuable indicators for the presence of greigite. The low \(\chi_{\text{ARM}} / \chi_{\text{SIRM}}\) (0.06–0.2 \(\times\) 10^{-3} mA^{-1}) values of the latter samples lie close to the range of authigenic inorganically precipitated greigite which is characterized by 0.058–0.084 \(\times\) 10^{-3} mA^{-1} (Snowball, 1997a), in contrast to high values observed for sediments containing bacterial magnetite (Snowball, 1994) and greigite (Reinholdsson et al., 2013, Fig. 3b). Only a few samples that did not acquire GRM lie in these latter areas, i.e., samples from the uppermost part of the core (cf. Fig. 2g, h). These results suggest that except for the latter samples, a significant contribution of magnetite and greigite magnetosomes can be ruled out as a source for rock-magnetic signals. Furthermore, we propose that greigite, within the GRM acquiring samples, formed out of a chemical process. However, this process was likely induced by biological activity (iron-reducing bacteria).

There appears to be a strong positive correlation between SIRM / \(\chi\) and \(\chi_{\text{ARM}} / \chi\) (Fig. 4c). The latter parameter is often used as a magnetic grain-size indicator for low-coercivity minerals, especially magnetite, while SIRM / \(\chi\) can be influenced by different mechanisms, ranging from changes in magnetic grain size (Peters and Dekkers, 2003) and magnetic mineralogy, including the presence of greigite (Frank et al., 2007; Fu et al., 2008; Nowaczyk et al., 2012; Reinholdsson et
al., 2013; Snowball, 1997b). Although paramagnetic minerals may contribute to the susceptibility, this bias would be expressed in both ratios, and a cross plot of those two parameters is valid to compare ferrimagnetic mineralogic properties between samples. The linear behavior of the two parameters indicates that SIRM / χ is influenced by magnetic grain-size effects. However, the bigger gradient for GRM samples implies that the presence of greigite is shifting SIRM to higher levels.

The MDF ARM is indicative of the hardness of magnetic minerals within the low-coercivity magnetic fraction (i.e. SD and PSD magnetite), and is therefore also affected by a preferential dissolution of fine magnetic particles. As outlined above, the S ratio signifies the relative concentration of high- vs. low-coercivity minerals and is influenced by relative increases in high-coercivity particles, or decreases in magnetite, the latter could also be induced by dissolution. From the MDF and S ratio plot, two clusters can be distinguished (Fig. 4d). Non-GRM samples have MDFs between 25 and 33 mT and a broad range of S ratios, whereas the MDF for GRM samples is higher than 30 mT and are mainly confined by S ratios higher than 0.96. The co-occurrence of high MDFs and GRM acquisition strongly suggests that increasing coercivity results from the presence of greigite. Moreover, the high S ratios reveal that greigite mostly contributes to the low-coercivity component of magnetization. For the non-GRM samples the pattern implies that relative increases of the high-coercivity fraction are not accompanied by changes within the low-coercivity mineral assemblage. A preferential dissolution of ferrimagnetic minerals, expressed as a drop in the S-Ratio, would be expected to be accompanied by changing coercivity also within low-coercivity assemblage (i.e., changing MDF). This is not observed for our samples. Moreover, the constant SIRM throughout unit 1, whereby HIRM shows variations (cf. Fig. 2c, d), rather indicates the addition of high-coercivity minerals, instead of a decrease in low-coercivity minerals. Further support for this hypothesis comes from the low TOC concentrations accompanying the low S ratio intervals. Enhanced magnetite dissolution is typically observed when TOC concentrations are elevated. The trend in the magnetic proxies, however, shows the opposite of what would be expected in the case of enhanced dissolution. The S ratio indicates even higher contents of low vs. high-coercivity minerals during interglacials with elevated TOC content. We therefore assume that, if reductive diagenesis had occurred, it affected the whole unit equally, and variation in the S ratio is indicative of increasing high-coercivity minerals within the detrital magnetic mineral fraction.

We also conducted high-temperature susceptibility measurements, in order to discriminate the iron sulfides. The susceptibility of all samples increased sharply above 400 °C and decreased above 500 °C (data not shown). The cooling curve has higher susceptibilities. This behavior is typical if reduced iron is oxidized to magnetite upon heating. This iron, however, can be derived from different mineral phases, e.g., from clay minerals (Bell et al., 1987) or iron sulfides and siderites.

### 5.2 Cluster analysis

To further evaluate temporal changes in the magnetic mineralogy of Lake Ohrid sediments we performed a cluster analysis. Based on the evaluation of the magnetic parameters, we included χ ARM / χ, χ ARM / SIRM, S ratio, MDF and ΔGRM / ΔNRM which are indicative of magnetic grain size, magnetic coercivity, and the occurrence of greigite.

Cluster center 3 is characterized by the lowest χ ARM / SIRM value, high S-Ratio, a high MDF of 40 mT and a high GRM (Table 2). These characteristics are often observed for sediments containing chemically produced greigite (Fig. 4); thus we infer that cluster 3 is related to greigite and other ferrimagnetic iron sulfides, e.g., pyrrhotite.

Cluster centers 1 and 2 have similarly low GRM and MDF, indicating the absence of greigite and undistinguish-

![Table 1. Magnetic properties of samples used for scanning electron microscopy.](http://www.biogeosciences.net/13/2093/2016/)

<table>
<thead>
<tr>
<th>Depth (m)</th>
<th>Age (kyr)</th>
<th>MDF (mT)</th>
<th>SIRM/χ (10⁻³ A m⁻¹)</th>
<th>ΔGRM/ΔNRM</th>
<th>χ ARM/SIRM (10⁻³ m A⁻¹)</th>
<th>Observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>10.53</td>
<td>26</td>
<td>27</td>
<td>9.8</td>
<td>0.00</td>
<td>0.24</td>
<td>Ti magnetite, magnetite Cr-Fe oxide, Mg-Cr spinel, Fe-Mn oxide</td>
</tr>
<tr>
<td>72.53</td>
<td>177</td>
<td>33</td>
<td>3.0</td>
<td>0.00</td>
<td>0.31</td>
<td>Ti magnetite, magnetite Cr-Fe oxide, Mg-Cr spinel, Fe-Mn oxide, siderite</td>
</tr>
<tr>
<td>117.83</td>
<td>270</td>
<td>36</td>
<td>11.6</td>
<td>0.00</td>
<td>0.12</td>
<td>(microcrystalline) Fe sulfide aggregates, idiomorphic greigite, siderite, Ti magnetite</td>
</tr>
<tr>
<td>153.83</td>
<td>360</td>
<td>45</td>
<td>29.1</td>
<td>0.01</td>
<td>0.09</td>
<td>fine-grained Fe-S (greigite?), Ti magnetite Cr-Fe oxide</td>
</tr>
<tr>
<td>158.63</td>
<td>378</td>
<td>47</td>
<td>22.5</td>
<td>0.70</td>
<td>0.10</td>
<td>fine-grained Fe-S (greigite?)</td>
</tr>
<tr>
<td>162.47</td>
<td>392</td>
<td>46</td>
<td>20.6</td>
<td>0.85</td>
<td>0.11</td>
<td>Ti magnetite fine-grained Fe-S nODULES</td>
</tr>
<tr>
<td>176.87</td>
<td>432</td>
<td>53</td>
<td>38.2</td>
<td>0.01</td>
<td>0.07</td>
<td>coarse-grained Fe-S nODULES, fine-grained greigite</td>
</tr>
</tbody>
</table>

www.biogeosciences.net/13/2093/2016/  
Biogeosciences, 13, 2093–2109, 2016
able different coercivities within the magnetically soft fraction. However, $\chi_{\text{ARM}} / \chi_s$, $\chi_{\text{ARM}} / \text{SIRM}$ and the $S$ ratio are lower in cluster 1 compared to cluster 2. As discussed in Sect. 5.1, the differences are due to the higher contribution of high-coercivity minerals, concurrent with a coarsening of the ferrimagnetic fraction.

The variations in cluster membership coefficients (Fig. 2a) captures the variability in magnetic proxies (Fig. 2b–j). Cluster 1 mainly relates to interglacial samples, whereas cluster 2 corresponds mainly to glacial samples in the upper part of the core (Fig. 5f). Glacials samples from the lower part of the core are associated with cluster 3.

Figure 3. SEM images of magnetic extracts; see also Table 1. (a) 72.53 m depth, detrital chromium iron oxides, titanomagnetite and magnetite with traces of Cr and Ti. (b) 10.53 m depth, idiomorphic and fragments of magnetites, traces of Ti and Cr. (c) 158.83 m depth, microcrystalline and framboidal iron sulfides within organic shell. (d) 162.47 m depth, chromium iron oxides, fine-grained greigite aggregates. (e) 153.83 m depth, microcrystalline iron sulfide nodule. (f) 153.83 m depth, microcrystalline iron sulfide aggregates, idiomorphic greigite crystals (arrows). (g) 117.83 m depth, coarse Fe-S nodules and fine-grained siderite grains (elongated particles). (h) 176.87 m depth, microcrystalline iron sulfide nodule.
Figure 4. Selected cross plots of magnetic proxy evaluation. Samples that acquire GRM (color code) cluster in different regions of the diagrams. (a) GRM samples are characterized by high $S$ ratios. A linear relationship between $\chi_{ARM}/SIRM$ and $S$ ratio relates to co-varying ferrimagnetic grain-size fining and increasing low- vs. high-coercivity magnetic mineral content. (b) GRM samples plot at $\chi_{ARM}/SIRM$ levels typical of authigenic, inorganically precipitated greigite. A few samples from the uppermost part of the core plot in the field of bacterial magnetite (dashed circle; Snowball, 1994) and greigite (green shaded area; Reinholdsson et al., 2013). (c) GRM samples have a distinctively different gradient compared to non-GRM samples in the SIRM/$\chi$ vs. $\chi_{ARM}/\chi$ plot. (d) GRM samples are characterized by high $S$ ratios and high $\text{MDF}_{ARM}$. Non-GRM samples show no relationship between the two parameters.

Table 2. Cluster center properties obtained from fuzzy c-means clustering.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>$\chi_{ARM}/\chi$ ($10^{-3}$ m A$^{-1}$)</th>
<th>$\chi_{ARM}/SIRM$ ($10^{-3}$ m A$^{-1}$)</th>
<th>S ratio</th>
<th>MDF$_{ARM}$ (%)</th>
<th>$\Delta GRM/\Delta NRM$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cluster 1</td>
<td>1.10</td>
<td>0.30</td>
<td>0.93</td>
<td>30.69</td>
<td>2.79</td>
</tr>
<tr>
<td>Cluster 2</td>
<td>5.53</td>
<td>0.55</td>
<td>0.97</td>
<td>29.85</td>
<td>2.08</td>
</tr>
<tr>
<td>Cluster 3</td>
<td>1.89</td>
<td>0.13</td>
<td>0.97</td>
<td>39.86</td>
<td>11.95</td>
</tr>
</tbody>
</table>

5.3 Iron sulfides in Lake Ohrid sediments

The concentration of magnetic minerals, approximated by remanence intensities (and susceptibilities), is relatively low in Lake Ohrid sediments in the upper unit. This gives a strong indication that the magnetic fraction was subject to reductive diagenesis. In unit 2, remanence intensities are enhanced, due to the presence of ferrimagnetic iron sulfides (Fig. 2). Concerning the occurrence of greigite and iron sulfides, the cluster analysis reveals two different patterns that modulate each other (Fig. 5f): firstly, the absence of iron sulfides in unit 1, and secondly, a cyclic occurrence of iron sulfides on glacial–interglacial timescales in unit 2. Besides iron sulfides, siderite, which also forms during (early) diagenesis, is contained in the sediments. When discussing the mechanism for diageneric iron-mineral formation, it is worthwhile also considering the occurrence of paramagnetic siderites.

5.4 Early vs. late diagenetic formation

For iron sulfides to precipitate, accessible Fe and S have to be available. In the course of early diagenesis sulfate is reduced during organic matter degradation. Typically, iron sulfides such as greigite and pyrite form in the sulfidic pore water zone, associated with the upward-migrating sulfate–methane transition zone (SMTZ), where $H_2S$ accumulates (Kasten et al., 1998).

In contrast, siderite mainly forms in the methanogenic zone, if pore waters have high CaCO$_3$ concentrations (e.g., Berner, 1981; Roberts, 2015). However, siderite and greigite (iron sulfides) can form at the same time if the rate of
iron reduction is higher than the rate of sulfate reduction (e.g. Pye, 1981, 1990). Sulfate concentrations are much lower in lakes (10–500 µM), compared to the ocean (28 mM; Holmer and Storkholm, 2001), and sulfate is depleted within the uppermost centimeter of the sediments. Thus, methanogenesis plays a more important role for organic matter degradation compared to marine environments (Capone and Kiene, 1988).

During late diagenesis, iron sulfides may precipitate from mineralized fluids (Sagnotti et al., 2010) if pore-water chemistry changes, i.e., by up- or downward-migrating dissolved H₂S species. Newly formed iron sulfides have been observed to overgrow earlier diagenetic iron minerals, such as siderite (Roberts and Weaver, 2005; Sagnotti et al., 2005).

From SEM observations, we find no indication that sulfides formed at a later stage than siderites. Sulfide nodules are much coarser than the apparently well-preserved siderite crystals (Fig. 3g). The shape of iron sulfide nodules rather appear to fill cavities of former organic compounds (Fig. 3c).

Siderite abundances are higher in the upper part of the core (Fig. 5e), where the ferrimagnetic iron sulfide proxies are low. In the older glacial of unit 2, siderite concentrations decrease when GRM, MDF_{ARM} and/or SIRM / \chi increase (Fig. 5b–d). This rather suggests that during early diagenesis one over the other, iron sulfide or siderite formation dominated.

Finally, this assumption is supported by unpublished relative paleointensity (RPI) data that will be presented elsewhere. If the ferrimagnetic iron sulfides formed later during diagenesis, a disruption of the match between the RPI trend from Lake Ohrid and global RPI stacks would be expected. However, the good quality of the RPI correlation suggests that the iron sulfide nodules, associated with elevated SIRM / \chi, grew during early diagenesis.

5.5 Large-scale shifts in redox conditions

The most prominent feature of the magnetic record is the transition from a dominance of iron sulfides in unit 2 (> 320 ka) and siderite in unit 1 (< 320 ka), suggesting a change in pore-water chemistry. Other proxy data sets from the Ohrid composite profile also indicate that lacustrine conditions changed at this boundary. As outlined above, siderite abundances are distinctly higher in the upper part of the core (Fig. 5e). Furthermore, interglacial TIC concentrations are generally lower in the upper unit 1 (up to 6 %) compared to unit 2 (up to 10 %, Fig. 5h), while interglacial TOC concentrations are relatively low in unit 2 (Fig. 2l). The higher concentrations of TIC relative to TOC were tentatively dis-
cussed to be dependent on higher ion concentrations in lake water due to increased evaporation (Lacey et al., 2016) during the deposition of the lower unit, but could also relate to enhanced organic matter degradation (Francck et al., 2015).

As outlined above, siderite typically forms during methanogenesis (Berner, 1981), which is the only process for organic matter degradation after sulfate is consumed. The shift of siderite to sulfide formation implies a shift in the redox conditions between unit 1 and 2. We propose that, during the deposition of unit 1, SO$_4^{2-}$ was rapidly consumed in the shallow sediments, or even within the bottom water. Magnetic iron oxides were readily reduced, but due to the lack of sufficient H$_2$S, siderite precipitated out of CaCO$_3$ supersaturated waters (Coleman et al., 1985), as was also observed in the sister lake Prespa (Leng et al., 2013). In contrast, the sulfidic zone apparently penetrated deeper into the sediments during the deposition of unit 2. Pyrite formation (Canfield and Berner, 1987) and, thus, precursors of pyrite, such as greigite, require sulfate concentrations > 1 mM. These prerequisites were apparently met; however, sulfate concentrations were probably still low, and not sufficient to transform metastable iron sulfides into pyrite.

The depth of the sulfidic zone is influenced by a number of different processes which can modulate each other. First of all, the input of sulfate or sulfide could differ. Generally, sulfate is mainly derived from sulfur-bearing weathered rocks in the catchment (Holmer and Storkholm, 2001). Enhanced erosion or a stronger chemical weathering could thus increase the sulfate supply to the lake. Sulfides may also derive from upward-migrating fluids in active tectonic settings. Secondly, even if sulfate fluxes are stable, sulfate concentrations increase when evaporation is enhanced. Another process of sulfate consumption in the sediments is linked to O$_2$ concentrations in the bottom water and to the accumulation and degradation of organic matter. Enhanced degradation of organic matter within the oxic zone, e.g., due to better ventilation of bottom water, would modify the oxidation state of bottom water and shift the sulfidic zone to a deeper depth.

At this point we cannot infer which process, or combination of processes, is responsible for the observed pattern. Given that Lake Ohrid is a subsiding basin, the water volume of the lake may have been smaller during deposition of unit 2. This could have led to a more regular deep mixing of Lake Ohrid and a slower sulfate consumption, both resulting in the sulfidic zone migrating deeper into the underlying sediments. This scenario is in line with higher TIC concentrations in the lower part of the core most likely due to higher ion concentrations and to comparably low TOC concentrations, linked to organic matter degradation. In addition, the glacial phases of low eccentricity, i.e., weaker seasonality (Fig. 5a). The annual temperature distribution may have had an additional effect on deep convection and lacustrine productivity of Lake Ohrid, which might have contributed to a change in mixing processes and the production and degradation of organic material.

5.6 Glacial–interglacial variability of iron sulfide formation

Ferrimagnetic iron sulfides (paramagnetic siderites) are contained within glacial sediments in the lower (upper) unit. Moreover, diagenetic Fe minerals, represented by the iron sulfide cluster (Fig. 5f) and siderite abundances (Fig. 5e), occur at peak Fe/Ca ratios and minimum TIC concentrations (Fig. 5) within interglacials.

In the older glacials (unit 2), siderite concentrations are high when GRM acquisition and SIRM/$\chi$ is relatively low, and vice versa. SIRM/$\chi$ rises, in turn, to maximum values when GRM and siderite content are reduced. Again, this highlights the importance of sulfate concentrations and Fe availability for the early diagenetic formation of iron minerals.

A further characterization of iron-mineral genesis in the course of redox conditions, however, needs to be further investigated by performing geochemical and mineralogical classification. Sulfur isotopes could further improve the understanding of the sulfur source, and a discrimination of the chemical composition of the iron sulfide nodules will help to understand the processes responsible for their formation and preservation.

5.7 High-coercivity minerals in Lake Ohrid sediments

Since the magnetic signal in unit 2 is dominated by the presence of diagenetic ferrimagnetic iron sulfides, we only investigated unit 1 (MIS 1–MIS 8) for changing lithogenic sediment supply (Fig. 5). Terrigenous input vs. limnic productivity is high during glacial, indicated by higher Fe/Ca ratio, and higher susceptibility (integrating ferri- and paramagnetic minerals) and low TIC concentrations. At the same time, the concentration of high-coercivity magnetic minerals increases within the magnetic fraction (indicated by the S ratio). Since this pattern is not due to preferential dissolution of magnetite (cf. Sect. 5.1), we propose that the composition of terrigenous input changed over glacial–interglacial timescales.

The catchment of Lake Ohrid comprises different lithologies (cf. Sect. 1) that are mirrored by the distribution of element concentrations in surface sediments (Vogel et al., 2010b). Vogel et al. (2010a) assumed that changes in Cr/Ti ratios on glacial–interglacial timescales result from either increased aeolian activity or a stronger erosion of soil material from sparsely vegetated soils. The detrital iron oxides in the magnetic extracts often contain Cr, which is a typical element for mantle rocks. The ophiolite belt, located west of the basin, is therefore a possible source for the magnetic minerals in Lake Ohrid. The magnetic mineralogy of the ophiolites (oceanic core complexes) in the Albanides consists mainly of magnetite (Maffione et al., 2014).

We observe a large similarity between the benthic $\delta^{18}$O stack (Lisiecki and Raymo, 2005) and the S ratio from Lake Ohrid (Fig. 3b, c). This suggests compositional changes
in the magnetic mineralogy reflect changing environmental conditions. During warm (and humid) interglacials, chemical weathering was enhanced and accumulation of soils and pedogenetic formation of (magnetic) minerals was promoted. However, as already proposed by Vogel et al. (2010a), vegetation cover prevented the erosion of the soil materials, and terrigenous magnetic minerals were diluted by biogenic sedimentary components. In the following glacial, vegetation cover decreased, as is indicated by arboreal pollen abundances from Lake Ohrid (Sadori et al., 2015) and soils were exposed and more susceptible to erosion. As a result, increased input of high-coercivity minerals, e.g., hematite and/or goethite, can be observed, the latter being the most widespread pedogenetic magnetic minerals (Cornell and Schwertmann, 2006; Vodyanitskii, 2010).

During interglacials, the $S$ ratio (Fig. 6d) and $\chi_{ARM}/SIRM$ (cf. Fig. 2g) show higher frequency variations, where low values approximate to local summer insolation minima (Fig. 6b). Offsets, e.g., during MIS 5, might be related to age uncertainties (±2000 years for the Lake Ohrid record; Francke et al., 2015). Changes in the magnetic mineralogy in concert with insolation are also mirrored by the cluster membership coefficients (Fig. 6a). Similar to the mechanism proposed above, these low-insolation phases correspond to relatively cold conditions and less dense vegetation cover, also visible in pollen abundance patterns (Sadori et al., 2015), thus increasing erosion of soil materials.

### 5.8 Bacterial magnetite in Lake Ohrid sediments?

In the uppermost 6 m of the core, covering the Holocene, $\chi_{ARM}/SIRM$ and SIRM reach the highest values of the sequence (Fig. 2). An increased input of lithogenic magnetic minerals relative to carbonates can be ruled out since TOC and TIC concentrations are high (Figs. 2 and 4).

One source for magnetic minerals, independent of detrital material supply, is the production of bacterial magnetite and greigite. Magnetotactic bacteria utilize dissolved iron that is available either in the water column or at the Fe$^{2+}$–Fe$^{3+}$ redox boundary in the sediment (Kopp and Kirschvink, 2008). These bacteria produce magnetite (Blakemore, 1975; Frankel et al., 1979) or greigite (Heywood et al., 1990; Mann et al., 1990) crystals, so-called magnetosomes, within or outside their cells. It was shown that production of bacterial magnetite is linked to increasing organic matter supply (Egli, 2004a; Roberts et al., 2011b; Snowball et al., 2002), at least for oxic lakes (Egli, 2004b). Moreover, the production of bacterial magnetite can be fostered by the input of nutri-
ents (Egli, 2004b). Magnetotactic bacteria producing greigite prefer reducing conditions, and greigite magnetosomes have a higher potential for preservation under sulfidic conditions (Chang et al., 2014; Vasiliev et al., 2008). Fine magnetite crystals have a potential for preservation if certain environmental conditions, e.g., supply of oxygen and concentration of hydrogen sulfide, are met (Canfield and Berner, 1987). In Fig. 3, the Holocene samples are the only ones that coincide with the area of bacterial magnetite and greigite. The concurrence of elevated TOC together with the fine-grained magnetic phase could therefore indicate the presence of bacterial magnetite. However, we cannot infer whether this occurrence is triggered by high TOC and/or nutrient input, or whether it results from a not-yet-completed dissolution of magnetosomes (Snowball, 1994).

6 Conclusions

Rock-magnetic data, in conjunction with sedimentological and geochemical data from the Lake Ohrid DEEP site, signify changing terrestrial climate conditions, as well as changes in the lacustrine system over the past 637 kyr. Magnetic parameters often associated with greigite are elevated in glacial periods in the lower part of the core (637–320 ka, unit 2). SEM investigations support the presence of greigite and/or other iron sulfides. Ferrimagnetic iron sulfides are absent in the upper part of the core (0–320 ka, unit 1), where instead siderite is abundant in glacial sediments. Since siderite typically forms during methanogenesis after SO₄ is consumed, we propose that a geochemical shift occurred in Lake Ohrid with higher (lower) sulfate availability during the deposition of the lower (upper) unit. Various mechanisms might be responsible for this pattern. However, based on higher TIC concentrations within the interglacial periods of the lower unit, which are probably linked to higher ion concentrations in the lake water, we suggest that sulfate flux was enhanced and/or sulfate was concentrated due to a smaller water volume or enhanced evaporation. Further studies on the iron sulfide mineralogy and sulfur isotopes are required to provide a better understanding of the sources of sulfur and processes responsible for differences in Fe-S morphology and chemistry.

The magnetic properties of sediments deposited during the past 320 kyr are also observed to signify changes in terrestrial environmental conditions on glacial and interglacial timescales. During glacial, high-coercivity magnetic minerals (e.g., hematite and goethite) that formed in the course of pedogenesis in preceding interglacials were deposited in the lake. In contrast, a rich catchment vegetation during interglacials limited the erosion of soil material and only minor detrital magnetite originating from physically weathered rocks was transported into Lake Ohrid. Millennial-scale variations in rock-magnetic properties, which are concurrent with changes in summer insolation, suggest that the proposed mechanism of vegetation expansion also influenced the erosion of soil materials on shorter timescales. Magnetic concentration parameters in the Holocene (upper 6 m) are enhanced, while carbonate and TOC concentrations, normally diluting the magnetic signal, are also high. Together with magnetic proxies for magnetic coercivity, these samples are suspected to contain bacterial magnetite. Overall, our findings demonstrate the valuable contribution of rock-magnetic methods to environmental studies, as they provide important information about a suite of different processes, comprising studies on terrestrial environmental conditions, sediment dynamics and internal lake processes.

Data availability

All rock-magnetic data are available from PANGAEA (Just et al., 2015) at doi:10.1594/PANGAEA.848639. Geochemical data are from Francke et al. (2016) and siderite data are from Lacey et al. (2016).

Acknowledgements. The SCOPSCO Lake Ohrid drilling campaign was funded by the ICDP, the German Ministry of Higher Education and Research, the German Research Foundation, the University of Cologne, the British Geological Survey, the INGV and CNR (both Italy), and the governments of the republics of Macedonia (FYROM) and Albania. Logistic support was provided by the Hydrobiological Institute in Ohrid. Drilling was carried out by Drilling, Observation and Sampling of the Earth’s Continental Crust (DOSECC) and using the Deep Lake Drilling System (DLDS). Special thanks to Beau Marshall and the drilling team. Ali Skinner and Martin Melles provided immense help and advice during logistic preparation and the drilling operation. We acknowledge many student assistants for sediment sampling. We also thank the two anonymous reviewers for constructive feedback on the manuscript. J. Just was financially supported through the Collaborative Research Centre 806, Deutsche Forschungsgemeinschaft.

Edited by: T. Wilke

References


www.biogeosciences.net/13/2093/2016/
J. Just et al.: Environmental control on the occurrence of high-coercivity magnetic minerals


Biogeosciences, 13, 2093–2109, 2016 www.biogeosciences.net/13/2093/2016/


Snowball, I. F.: The detection of single-domain greigite (Fe3S4) using rotational remanent magnetization (RRM) and the effective gyro field (Bg): mineral magnetic and palaeomagnetic applications, Geophys. J. Int., 130, 704–716, 1997a.


