

Contents lists available at ScienceDirect

Physics of the Earth and Planetary Interiors

journal homepage: www.elsevier.com/locate/pepi

Diagenetic alteration of natural Fe–Ti oxides identified by energy dispersive spectroscopy and low-temperature magnetic remanence and hysteresis measurements

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ARTICLE INFO

Article history: Received 20 December 2007 Received in revised form 24 July 2008 Accepted 6 August 2008

Keywords: Magnetic Fe-Ti oxides Energy dispersive spectroscopy (EDS) Marine sediments Scanning electron microscopy (SEM) Low-temperature magnetism Early diagenesis

ABSTRACT

Low-temperature (LT) magnetic remanence and hysteresis measurements, in the range 300–5 K, were combined with energy dispersive spectroscopy (EDS) in order to characterize the magnetic inventory of strongly diagenetically altered sediments originating from the Niger deep-sea fan. We demonstrate the possibility of distinguishing between different compositions of members of the magnetite–ulvöspinel and ilmenite–hematite solid solution series on a set of five representative samples, two from the upper suboxic and three from the lower sulfidic anoxic zone of gravity core GeoB 4901. Highly sensitive LT magnetic measurements were performed on magnetic extracts resulting in large differences in the magnetic behavior between samples from the different layers. This emphasizes that both Fe–Ti oxide phases occur in different proportions in the two geochemical environments.

Most prominent are variations in the coercivity sensitive parameter coercive field (B_C). At roomtemperature (RT) hysteresis loops for all extracts are narrow and yield low coercivities (6–13 mT). With decreasing temperature the loops become more pronounced and wider. At 5 K an approximately 5-fold increase in B_C for the suboxic samples contrasts a 20–25-fold increase for the samples from the anoxic zone. We demonstrate that this distinct increase in B_C at LT corresponds to the increasing proportion of the Ti-rich hemoilmenite phase, while Fe-rich (titano-)magnetite dominates the magnetic signal at RT. This trend is also seen in the room-temperature saturation isothermal remanent magnetization (RT-SIRM) cycles: suboxic samples show remanence curves dominated by Fe-rich mineral phases while anoxic samples display curves clearly dominated by Ti-rich particles.

We show that the EDS intensity ratios of the characteristic Fe K_{α} and Ti K_{α} lines of the Fe–Ti oxides may be used to differentiate between members of the magnetite–ulvöspinel and ilmenite–hematite solid solution series. Furthermore it is possible to calculate an approximate composition for each grain if the intensity ratios of natural particles are linked to well-known standards. Thus, element spectra with high Fe/Ti intensity ratios were found to be rather typical of titanomagnetite while low Fe/Ti ratios are indicative of hemoilmenite.

The EDS analyses confirm the LT magnetic results, Fe-rich magnetic phases dominate in the upper suboxic environment whereas Ti-rich magnetic phases comprise the majority of particles in the lower anoxic domain: The mineral assemblage of the upper suboxic environments is composed of magnetite (\sim 19%), titanomagnetite (\sim 62%), hemoilmenite (\sim 17%) and \sim 2% other particles. In the lower anoxic sediments, reductive diagenetic alteration has resulted in more extensive depletion of the (titano-)magnetite phase, resulting in a relative enrichment of the hemoilmenite phase (\sim 66%). In these strongly anoxic sediments stoichiometric magnetite is barely preserved and only \sim 5% titanomagnetite was detected. The remaining \sim 28% comprises Ti-rich particles such as pseudobrookite or rutile.

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1. Introduction

A detailed rock magnetic study of diagenetically altered sediments from the Niger deep-sea fan (GeoB 4901) was reported recently by Dillon and Bleil (2006). Room-temperature hysteresis, IRM, ARM, high- and low-temperature data were combined

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^{0031-9201/\$ –} see front matter @ 2008 Elsevier B.V. All rights reserved. doi:10.1016/j.pepi.2008.08.003



Fig. 1. Location of sediment coring site GeoB 4901 in 2184 m water depth on the southeastern flank of Niger deep-sea fan (02°40.7'N/06°43.2'E). Isobaths are shown at 1000 m intervals according to Intergovernmental Oceanographic Commission (1994).

to provide a comprehensive interpretation of the magnetic assemblage and its modification by a variety of diagenetic processes. The authors suggested that the vast majority of the magnetic assemblage reaching the core location was composed of eroded volcanic material originating from the Cameroon Volcanic Line. The volcanic assemblage was considered to be of variable composition, containing Ti-poor and Ti-rich titanomagnetites and ilmenohematites/hemoilmenites (Herrero-Bervera et al., 2004; Ubangoh et al., 2005).

At gravity core location GeoB 4901 (02°40.7'N/06°43.2'E, 2184 m water depth, Schulz et al., 1998; Fig. 1) the degradation of organic matter results in a specific redox zonation of the sediment column, with only the upper 10 cm being oxic. Suboxic conditions are characterized by dissolved manganese in the pore water and are established down to 12.50 m. Anoxic conditions prevail below this level where anaerobic oxidation of methane results in free H₂S in the pore water (Heuer, 2003). In this distinct redox zonation the magnetic iron oxide mineral inventory undergoes a two-stage diagenetic alteration. The modern iron redox boundary (\sim 2.95 m) is characterized by a drastic decline in magnetic mineral content, a coarsening of the grain-size spectrum, and a reduction in coercivity at room temperature. Such reductive diagenesis in the suboxic zone is a common and frequently studied phenomenon in organic-rich marine sediments (e.g. Karlin and Levi, 1985; Canfield and Berner, 1987). Below the second diagenetic horizon, the suboxic/sulfidic anoxic transition (\sim 12.50 m), a further but less pronounced decrease in magnetic concentration and grain size occurs. The contribution of finer magnetic particles and the proportion of higher coercive magnetic constituents substantially increase in the anoxic environment (Dillon and Bleil, 2006).

Diagenetic alteration of the primary detrital magnetic mineral inventory was found to be dependent on both particle size and Ti-content, with the smallest particles dissolving first and the Tirich component being more resistant to the reducing conditions. Possible candidates for the preserved grains are titanomagnetite

 $(Fe_{3-x}Ti_xO_4)$ and hemoilmenite $(Fe_{2-v}Ti_vO_3)$ with compositions of x > 0.75 and y > 0.75, respectively. Both mineral phases with Ticontents ≥ 0.75 are paramagnetic at ambient temperatures and exhibit high magnetic stabilities at low-temperature (Bozorth et al., 1957; Schmidbauer and Readman, 1982; Ishikawa et al., 1985). More recent work by Burton et al. (2008) states that this is actually the case for hemoilmenite compositions of v > 0.8. Nevertheless. a definite distinction between the two mineral phases remains unresolved. Titanomagnetites with such compositions are rather uncommon as natural compounds. Therefore it appears more conceivable that Fe–Ti oxides of y > 0.75 constitutes hemoilmenite. The role of members of the ilmenite-hematite solid solution series in a magnetic mineral assemblage is often disregarded, although these particles exist in a large range of compositions and are frequent constituents of many types of rocks. However, the presence of (titano-)magnetite usually masks the signature of the magnetic members of the ilmenite-hematite solid solution series when both are present in a sample because the saturation moment of magnetite is at least three times greater than that of e.g. hemoilmenite (y=0.63) at room temperature (Nord and Lawson, 1989). For the sake of clarity in nomenclature we will use the term 'hemoilmenite' for compositions of $y \ge 0.5$ and the term 'ilmenohematite' for mineral phases with y < 0.5. Detailed compositions are expressed using the definitions of Nord and Lawson (1989), e.g. Ilm80Hem20 for y = 0.8.

At room-temperature members of the ilmenite–hematite solid solution series are either para-, ferri-, or antiferromagnetic, depending on their individual cation ordering and spin structure (e.g. McEnroe et al., 2000; Burton et al., 2008). Ilmenohematites with compositions of 0 < y < 0.5 are antiferromagnetic at room temperature and their magnetic moments are aligned as in hematite (Warner et al., 1972). Ferrimagnetic ordering in hemoilmenite occurs for intermediate compositions (0.5 < y < 0.8) and is related to the ordering of Fe and Ti on separate sub-lattices (Ishikawa, 1985; Burton et al., 2008). Compositions with y > 0.8 are paramagnetic

at room temperature, but magnetic ordering does occur at lower temperatures, e.g. the end-member ilmenite is antiferromagnetic below its Néel temperature of \sim 58 K (Kato et al., 1982).

This study has two aims: first to identify the low-temperature remanence and hysteresis characteristics that will help to discriminate between the members of the magnetite-ulvöspinel and ilmenite-hematite solid solution series. This will expand the lowtemperature database and therefore our understanding of natural samples. Secondly, the applied low-temperature magnetic measurements provide an insight into the environmental implications for the gravity core retrieved from the Niger deep-sea fan, by demonstrating the compositional changes of the magnetic mineral assemblage during progressive sub- and anoxic diagenesis. The rock magnetic results are linked to findings of quantitative SEM energy dispersive spectroscopy (EDS) analyses. In particular, the intensity ratios of the characteristic Fe K_{α} and Ti K_{α} lines are used to differentiate between the mineral phases of the magnetite-ulvöspinel and ilmenite-hematite solid solution series. This combination of magnetic and electron microscopic techniques is helpful in understanding the post-depositional processes in the sedimentary setting of the Niger deep-sea fan and therefore allows us to draw conclusions on the alteration of the primary continental input.

2. Materials and methods

2.1. Sample selection and treatment

Magnetic extraction following the method of Petersen et al. (1986) was applied to obtain magnetic concentrates for five representative samples along the depth profile of gravity core GeoB 4901 from (a) the upper suboxic sediments above the Fe-redox boundary (1.65 m), (b) the upper suboxic sediments below the Feredox boundary (2.95 m), and (c) to (e) from the anoxic sulfidic sediments (13.25, 14.55 and 15.25 m; see Fig. 2). The enrichment of the magnetic fraction was necessary because the starting concentration is too low for meaningful bulk sediment measurements. Mass determination of the extracts is not practical here, since the typical amount for these magnetic extracts is below the precision of a microbalance ($\sigma = 10^{-5}$ g). The sample from 1.65 m depth represents the zone where diagenetic influences are minimal and hence characterizes the primary magnetic mineralogy. The main interest of sampling at 2.95 m was to detect the potential existence of biogenic magnetite, which may form around the modern Fe-redox boundary (Hilgenfeldt, 2000). Samples between 13.25 and 15.25 m were selected to characterize the magnetic mineral assemblage in the anoxic sediments (Fig. 2).

2.2. Scanning electron microscopy

SEM analysis included backscatter electron (BSE) imaging (Lloyd, 1985) performed on the magnetic extracts of selected unconsolidated samples from depths of 1.65, 14.55 and 15.25 m depth, using a *FEI XL30 SFEG* scanning electron microscope at the EMU (Electron Microscopy Utrecht, The Netherlands). The SEM was operated at an acceleration voltage between 12 and 30 kV, with a \sim 2 nA beam current and a spot size of \sim 1.5 nm. Sample preparation for SEM analysis was carried out as described in Franke et al. (2007a). The elemental composition was acquired using energy X-ray dispersive spectroscopy (EDS; Goldstein et al., 1992). The obtained element spectra were (semi-)quantified using the '*Remote SEM Quant Phiroz*' program version 3.4. All element spectra are normalized to the height of their oxygen peak. Background noise was subtracted before calculation of the element composition. The SEM



Fig. 2. (a) Depth profile of the volume specific bulk susceptibility κ delineating variations in the concentration of magnetic minerals. (b) Pore water chemistry at Site GeoB 4901 (Heuer, 2003). All parameters are plotted against a linear depth scale and a non-linear age scale (Adegbie, 2001) including marine oxygen isotope stages (MIS) and terminations (T). Grey shading indicates cold periods. Horizontal hatched bars mark the modern iron redox boundary (~2.85 m) and the suboxic/sulfidic anoxic transition (~12.5 m). Arrows at the κ profile denote the horizons, where detailed magnetic analyses have been performed.

analyses are restricted to the resolvable μ m-particle fraction of the extracts and EDS was performed for about 40–50 particles per sample. The upper suboxic environment is represented by the sample from 1.65 m depth. Samples from 14.55 and 15.25 m depth represent the anoxic zone. In the following the SEM results of the two samples from 14.55 and 15.25 m depth are combined and will be referred to as anoxic samples.

2.3. Low-temperature magnetic measurements

Low-temperature magnetic measurements were performed on a superconducting *Quantum Design* Magnetic Property Measurement System (MPMS), with an instrumental noise level of $\sim 10^{-11}$ Am². Two types of low-temperature measurements were performed in this study: (1) saturation isothermal remanent magnetization imparted at room temperature (RT-SIRM) by applying a maximum field of 5 T at 300 K followed by continuous cooling in zero field to 5 K and subsequent continuous warming back to 300 K. (2) Hysteresis loops measured between magnetic peak fields of \pm 5 T at 15 distinct temperatures between 5 and 295 K. From these loops the temperature dependence of the standard hysteresis parameters such as saturation magnetization (M_S), saturation remanent magnetization (M_{RS}) and coercive force (B_C) were determined.

3. Scanning electron microscopic results

3.1. Discrimination between titanomagnetite and hemoilmenite

Studies such as Franke et al. (2007b) and Lattard et al. (2005) have shown that the EDS intensity ratios of the characteristic Fe K_{α} and Ti K_{α} lines can be used under certain conditions to differentiate between mineral phases of the magnetite-ulvöspinel and ilmenite-hematite solid solution series. Element spectra with high Fe/Ti intensity ratios were found to be rather typical of titanomagnetite while low Fe/Ti ratios are indicative of hemoilmenite. With increasing Ti-content the Fe/Ti ratio decreases for both solid solution series, the lowest possible value is Fe/Ti = 2for the magnetite-ulvöspinel solid solution series and Fe/Ti=1 for the ilmenite-hematite solid solution series according to their respective stoichiometric formulas. Ratios of Fe/Ti < 1 constitute members of the ternary system between rutile (TiO₂) and the pseudobrookite-ferropseudobrookite solid solution series (Fe₂TiO₅-FeTi₂O₅). Therefore particles with ratios of Fe/Ti > 2 could be theoretically either members of the magnetite-ulvöspinel or the ilmenite-hematite solid solution series, while particles with ratios of $1 \le \text{Fe}/\text{Ti} \le 2$ can only be members of the ilmenite-hematite solid solution series.

To test this method, EDS spectra were taken from a defined synthetic sample (6F72x2.4) of well-known composition, which was provided by the Mineralogical Institute of the University of Heidelberg. The sample was synthesized at 1300 °C in the Fe–Ti–O system and contains titanomagnetite (TM63) and synthetic hemoilmenite (=ferrian ilmenite; Ilm76Hem24). This sample was subsequently used as a pre-defined standard to which the EDS peak intensities of the unconsolidated natural particles of this study were compared. Another advantage of the synthetic sample is its ideal flat polished surface, which means that considerable scattering effects can be neglected.

The element spectra were derived on the same SEM instrument for both the synthetic and natural samples. Fig. 3a shows a backscatter electron micrograph of the synthetic sample 6F72x2.4. The grey tones in the BSE image reflect the different chemical compositions of the two mineral phases, with lighter areas corresponding to titanomagnetite (Fe-richer areas) and darker areas corresponding to members of the ilmenite-hematite solid solution series (Fe-poorer areas). Completely black or white spots mark pores in the polished sample, where surface charging might have an effect due to poor carbon coating. The two mineral phases can be discriminated by their EDS element spectra (Fig. 3b and c), since the intensity ratios of the characteristic Fe K_{α} and Ti K_{α} lines are distinct. In Table 1 the semi-quantified Fe and Ti contents are listed for ten spot checks analyzed on the polished section of the synthetic sample. The values were used to calculate the Fe/Ti ratios and the resulting titanomagnetite (TM) and hemoilmenite (IlmHem) mean compositions. Spectra from TM (lighter area) show Fe/Ti ratios >2, corresponding to quantitative EDS spectra for which the Fe-peak is equal or higher than the Ti-peak. Spectra from hemoilmenite (darker areas) have ratios $1 \le Fe/Ti \le 2$, showing Ti-peaks that are higher than the Fe-peaks. This can be easily explained by the position of the Ti K_{α} (at 5.896 eV) and Fe K_{α} (at 6.403 eV) lines in the spectra, e.g. a particle with the same peak height for both elements contains therefore less Ti than Fe. The average composition of the synthetic sample 6F72x2.4 was calculated as TM68 and Ilm78Hem22. These values are only slightly higher than the compositions (TM63 and Ilm76Hem24) derived from published electron microprobe (EMP) analyses and inductively coupled plasma atomic emission spectrometry (ICP-AES) reported by Lattard et al. (2005, 2006). The error of our EDS method is therefore less than 3%. This discrepancy is presumably due to slightly inhomogeneous chemical compositions within the sample from the rim to the center of the synthesized pellet (Engelmann, personal communication, 2006). These results underline that it is in principle possible to discriminate between titanomagnetite and hemoilmenite phases on the basis of the element intensity ratio Fe/Ti, hence that it is possible to calculate a close approximate composition for each individual particle.

3.2. Magnetic particle discrimination

The selected beam strength between 12 and 30 kV used for the quantitative EDS analysis detects all main elements within the Fe–Ti–O system and furthermore all potential minor element contributions which might be present in the mineral grains. The elements V, Cr, Al and Ti are common substitutions in hematite, whereas Mg and Mn are rather typical constituents in ilmenite. Besides Ti, magnetite may also contain the elements Mg, Mn, Cr and Al.

In practice, the analyzed particles from both solid solution series mainly show substitutions of Mg and Al. Other minor element contributions were rarely detected in our samples and were found in less than 20% of all analyzed particles. The contribution of these additional minor elements to the individual EDS spectra is usually less than 3%. Therefore the effect of minor elements other than Al and Mg can be neglected for this study. This concurs with the fact that Mg and Al are the most common cation impurities in natural titanomagnetites and hemoilmenites found in igneous rocks (Creer et al., 1975).

Fig. 4a and b show the cation element content in at% for the main elements Fe and Ti and the minor abundant elements Mg and Al for all examined particles, divided into suboxic and anoxic environments. To generally consider the effect of substitution of Fe with either Mg or Al, regardless for which member of both solid solution series, the total of the three elements Fe, Mg and Al was calculated and will be referred to as Fe_{Σ} (=Fe+Al+Mg) in the following interpretations.

The magnetic mineral assemblage of the upper suboxic environment is dominated by Fe-rich mineral phases with fairly low Ti-contents (Fig. 4a). In contrast, the Ti-content increases notably and even exceeds the Fe-content in the particles of the lower anoxic samples (Fig. 4b). In Fig. 4c and Fig. 4d representative element spectra of typical particles from the magnetic extracts are shown, originating from the upper suboxic and lower anoxic domain. These EDS analyses clearly show that the natural magnetic grains contain minor amounts of other ions than Fe and Ti. The presence of the supplementary elements, such as Na, Mg, Al and Si support the assumption of the detrital origin of these particles. Small amounts of Al and Si are most likely also due to clay mineral coatings on the grains.

In Table 1 the element ratios of Fe_{Σ}/Ti for each analyzed particle are divided into the $Fe_{\Sigma}/Ti \ge 2$; $1 \le Fe_{\Sigma}/Ti < 2$; $0.5 \le Fe_{\Sigma}/Ti < 1$ and $Fe_{\Sigma}/Ti < 0.5$ fractions. In the upper suboxic zone ~19% of the examined particles exhibit spectra without any Ti-content, comparable to spectrum (1) in Fig. 4c. They represent detrital magnetite, a common component in the upper suboxic core section. However, the majority of the magnetic particles contain Ti, with an element ratio Fe_{Σ}/Ti varying between 0.75 and 13.89; 79% of these particles have Fe_{Σ}/Ti ratios ≥ 2 , only 2% have Fe_{Σ}/Ti ratios <1 (Table 2).

According to the (semi-)quantitative EDS spectra, the Ti-content for particles with ratios of $Fe_{\Sigma}/Ti \ge 2$ varies between 0.4 and 4.9 at%. As already explained in the precedent section, this could



Fig. 3. (a) Backscattered electron micrograph of synthetic sample 6F72x2.4, light grey areas correspond to TM63 (Fe-richer) and darker grey areas correspond to Ilm76Hem24 (Fe-poorer). (b) EDS spectrum of the TM63 phase from the spot marked in (a). (c) EDS spectrum of the Ilm76Hem24 phase from the spot marked in (a).

 Table 1

 Cation element content in at% for the elements Fe and Ti of the synthetic sample 6F72x2.4

Sample name	Spot	Mineral phase	Ti [at%]	Fe [at%]	Fe/Ti	TM calculated	IlmHem calculated
6F72x2.4	Lighter area	TM	4.70	17.51	3.73	63	
6F72x2.4	Darker area	IlmHem	8.53	12.67	1.49		80/20
6F72x2.4	Lighter area	TM	4.75	16.51	3.48	67	
6F72x2.4	Darker area	IlmHem	8.06	12.53	1.55		78/22
6F72x2.4	Darker area	IlmHem	7.96	12.62	1.59		77/23
6F72x2.4	Lighter area	TM	4.83	16.25	3.36	69	
6F72x2.4	Lighter area	TM	5.20	17.53	3.37	69	
6F72x2.4	Lighter area	TM	5.23	18.57	3.55	66	
6F72x2.4	Darker area	IlmHem	7.96	12.73	1.60		77/23
6F72x2.4	Lighter area	TM	4.81	17.28	3.59	65	



Fig. 4. Univariate box-and-whisker plots of the cation element content for the main elements Fe and Ti and minor contributions of Mg and Al of magnetic extracts from (a) the suboxic and (b) the anoxic sediments. Fe_{Σ} equals the sum of the three elements Fe, Mg and Al. The boxes represent the median values and the interquartile range, the whiskers mark the total data range. Symbols (+) display outlier samples from the respective data group; (c) and (d) show typical element spectra for particles representing the different Fe–Ti oxide phases detected in the two geochemical environments (for further explanations see text).

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$\Sigma/Ti < 0.5$ Fe Σ/T	i<1 Fe _Σ /7	Γi<2 Fe _Σ /Ti	≥ 2 Ti/Fe _{Σ} = 0
2%	-	79%	19%
-	-	100%	-
26%	47%	24%	-
8%	-	92%	-
5	z/Ti < 0.5 Fe _Σ /T 2% - 26% 8%	$r/Ti < 0.5$ Fe _{Σ} /Ti < 1 Fe _{Σ} /Ti 2% - - - 26% 47% 8% -	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

Table 2Distribution of the Fe_{Σ}/Ti ratio for N particles examined

either correspond to a titanomagnetite composition between TM20 and TM90 or to compositions between Ilm13Hem87 and Ilm60Hem40 for members of the ilmenite–hematite solid solution series. For $0 \ge x > 0.75$ these assumed titanomagnetite compositions are conceivable. However, titanomagnetites with x > 0.75 are quite uncommon in nature (Petersen and Bleil, 1982). Instead it is more likely that grains with x > 0.75 represent hemoilmenite particles instead, which then would correspond to compositions between Ilm50Hem50 and Ilm60Hem40. Hemoilmenites of such compositions are typical for volcanogenic rocks that underwent high-temperature deuteric oxidation processes.

We have to keep in mind however, that natural mineral grains of the magnetite–ulvöspinel and ilmenite–hematite solid solution series occur rarely in completely homogenous compositions. In fact paragenesis between Ti-poor (titano-)magnetite or ilmeno-hematite and Ti-rich hemoilmenite is a common aftereffect of oxy-exsolution of volcanic rocks. Depending on the progression of such alterations, the crystal matrix might contain exsolution lamellae in nano- or even micrometer scale (e.g. McEnroe et al., 2001; Garming et al., 2007). The EDS analyses has a spatial resolution of ~0.5 μ m, therefore it was possible to resolve inhomogeneities up to that limit. Beyond this spatial resolution we might have to deal to a certain degree with overlap analyses, if exsolution features are present in a given grain. For particles larger than 0.5 μ m, multiple spot analyses were performed to check local inhomogeneities or to derive average compositions.

Additionally, magnetic inclusions in silicates were observed. They make up ~5% of the magnetic assemblage in the upper suboxic zone. The major elements in the measured spectra are Si and O, which are accompanied by Fe and Ti as minor components. Calculated element compositions for the magnetic inclusions show Fe_{Σ}/Ti ratios of \geq 2, resulting in either TM15 to TM60 or alternatively Ilm09Hem91 to Ilm40Hem60. Such typical Ti-poor composites have been found in various sedimentary environments as described by e.g. Garming et al. (2005).

Shrinking cracks are evidence for a mild maghemitization (Dunlop and Özdemir, 1997), but only a small number was observed in the upper suboxic sample. This suggests that (titano-)maghemite is of low importance in these samples. In addition to the observed microparticles, a considerable number of much smaller grains in the nm-range are present in the sample from the upper suboxic zone. These grains are beyond the spatial resolution of the used SEM. This is indicative for the presence of small PSD/SD particles.

In contrast to the analysis of the suboxic sample, only minor amounts of nm-sized particles were observed in the magnetic extracts from 14.55 and 15.25 m. These samples mainly consist of somewhat "coarser" grains in the grain-size range from ~2 to 5 μ m. Non-magnetic particles, such as silicates and feldspars are more abundant here than in the upper suboxic sample. This has to be kept in mind for the interpretation of the hysteresis data.

The average Ti-content of all particles examined is notably higher for the anoxic samples (\sim 4.06 at%) compared to the suboxic samples (\sim 1.61 at%) and even exceeds the meager Fe-content (Fig. 4b). When taking the Mg and Al concentrations into account, the mean value of Fe Σ is slightly higher than the average Ti-content.

In terms of numbers of grains the Ti-rich mineral phases clearly dominate the (magnetic) mineral inventory in the lower anoxic sediments. Here the ratios Fe_{Σ}/Ti vary between 0.21 and 4.87, particles without any Ti-content were no longer observed. Only 24% of the examined grains exhibit Fe_{Σ}/Ti ratios ≥ 2 . These particles would either refer to compositions between TM51 and TM99 or to compositions of Ilm34Hem66 to Ilm66Hem34. Particles with $1 \leq Fe_{\Sigma}/Ti < 2$ (47%) appear to be the most abundant members of the ilmenite–hematite solid solution series. Their Ti-content varies between 0.95 and 7.04 at%, corresponding to Ilm68Hem32 and Ilm99Hem01, respectively.

Additionally, a considerable number of particles yield Fe_{Σ}/Ti ratios of <1 (26%) or even <0.5 (3%). The element spectra for this particle type exhibit much higher Ti- than Fe-content (Fig. 4; compare spectrum 5). This composition corresponds rather to the pseudobrookite-ferropseudobrookite solid solution series. Pseudobrookite may be present in the detrital Fe–Ti oxide assemblage as minor constituents within titanomagnetite and –hematite grains as a result of high-temperature oxidation during the initial cooling of igneous rocks (Reynolds and Goldhaber, 1978; Frost and Lindsley, 1991). Another possibility for the genesis of such Tirich mineral phases is the further alteration of members of the ilmenite–hematite solid solution series towards hematite with fine rims of anatase (Houston and Murphy, 1962). This would enrich the elemental bulk composition of the grains in terms of their TiO₂ content.

The number of silicates with magnetic inclusions was found to be reasonably higher (14%) in the anoxic sediments compared to the suboxic sample (5%). The majority of these inclusions (92%) have Fe_{Σ}/Ti ratios \geq 2 (resulting in compositions of TM09 to TM75 or Ilm06Hem94 to Ilm50Hem50), only 8% of the inclusions show Fe_{Σ}/Ti ratios of <1.

3.3. Summary of SEM results

As mentioned above, titanomagnetites with x > 0.75 are quite rare in nature (Petersen and Bleil, 1982). The range of the composition for hemoilmenite strongly relates to the bulk chemistry of the rock, but lies typically above y = 0.5 (Petersen and Bleil, 1973). Considering these restrictions, the approximate abundances of the various Fe–Ti oxides in the magnetic mineral assemblage of the investigated samples can be estimated (Table 3).

Table 3

Relative abundances of mineral components in the upper suboxic and lower anoxic magnetic extracts

Component	Suboxic [%]	Anoxic [%]
Magnetite	18.9	0
Titanomagnetite	62.3	5.4
Hemoilmenite	16.9	66.2
Others	1.9	28.4

The mineral assemblage of the upper suboxic environment constitutes ~19% magnetite, ~62% titanomagnetite (TM20 to TM70), ~17% hemoilmenite (Ilm50Hem50 to Ilm60Hem40) and ~2% other particles. The composition of magnetic inclusions in the siliceous matrix varies between TM15 and TM60. The main magnetic mineral component in the magnetic assemblage of the lower suboxic environment is hemoilmenite with an abundance of ~66% and compositions varying between Ilm53Hem47 and Ilm99Hem01. Pure magnetite is no longer present and only ~5% titanomagnetite was found. The remaining ~28% are Ti-rich particles, such as (ferro-) pseudobrookite or rutile. When titanomagnetite is preserved as inclusions in a siliceous matrix it apparently withstands diagenetic alteration even in the anoxic environment.

4. Low-temperature magnetic results

4.1. Room-temperature SIRM (RT-SIRM)

Fig. 5 shows the RT-SIRM curves normalized to their initial value at 300K and the respective derivatives for all analyzed samples. The samples from the suboxic zone (Fig. 5a and b) clearly show ferrimagnetic dominated curves typical of (titano-)magnetite as described by Özdemir and Dunlop (2003) and Kosterov (2003). The RT-SIRM first increases upon cooling, reaching a maximum around 220 K (Fig. 5a) and 230 K (Fig. 5b). According to Dankers (1978) such an increase upon cooling is related to an increase in the spontaneous magnetization of titanomagnetite phases whose Curie point is above the temperature where the remanence is acquired. For such cases the spontaneous magnetization has not reached its maximum intensity at 300 K, which results in an increase while cooling. Upon further cooling the RT-SIRM curves decrease with decreasing temperature. At 5 K some 83% (Fig. 5a) and 69% (Fig. 5b) of the initial RT-SIRM are demagnetized. At 100 to 110 K the maximum gradient of the cooling curve is reached, corresponding to the temperature interval that is indicative for the Verwey structural phase transition of magnetite. This complies with results from zero field cooling (ZFC) and field cooling (FC) thermal demagnetization experiments, published by Dillon and Bleil (2006), which revealed partially suppressed Verwey transitions in the same temperature interval. During warming from 5 to 300 K the RT-SIRM retraces the cooling curve between 5 and 50 K. Upon further warming only limited remanence recovery is observed when passing back through the Verwey transition. At 300 K about 87% (Fig. 5a) and 80% (Fig. 5b) of the initial RT-SIRM is recovered.

The RT-SIRM behavior of samples from the anoxic zone differs strongly to that of the samples from the suboxic layers. However, within the anoxic zone all samples (Fig. 5c-e) reveal very similar low-temperature behavior with comparable remanence memory. The RT-SIRM cooling curves show a continuous gradual increase between 300 and 5 K, where the RT-SIRM is 46 to 69% higher than the initial remanence. Upon warming the curves resemble the corresponding cooling curves back to \sim 100 K. However, above 100 K the warming curves are below the cooling curves. The remanence loss when returning to 300 K accounts for 86-89% of the initial RT-SIRM. No specific magnetic transitions were observed on warming, though the first derivatives of samples 13.25 and 15.25 m display an inflection around 210K. This may be indicative of the transition of hemoilmenite with an approximate composition of y = 0.85from ferrimagnetism/superparamagnetism back to paramagnetism (Burton et al., 2008).

4.2. Low-temperature magnetic hysteresis

For all five magnetic extracts the temperature dependence of their hysteresis loops was examined between 5 and 295 K.



Fig. 5. Temperature variation of RT-SIRM during zero field cooling from 300 to 5 K (black solid line) and zero field warming back to 300 K (grey solid line). First derivative curves of the RT-SIRM curves are shown as dashed lines with the same color code.



Fig. 6. Uncorrected (black solid line), linear slope corrected (grey solid line) and paramagnetic slope corrected (grey dashed line) hysteresis data shown for samples from (a) 1.65 m and (b) 15.25 m core depth measured at 5 K. Filled triangles indicate the high field slopes determined by least squares fit of data measured between 4 and 5 T; filled diamonds depict the paramagnetic slopes derived from the equation $m(B, T) = a \cdot \tanh(B/b)$.

The measured hysteresis loops contain diamagnetic contributions originating from the sample holder (gelatine capsule) and non-ferromagnetic sediment constituents in the extracts such as feldspars and silicates, which to a certain extent were included in the extract. The temperature dependence of the paramagnetic contribution was calculated, using the relationship $m(B, T) = a \cdot \tanh(B/b)$, where m(B, T) equals the measured magnetic moment in the applied field *B* at the temperature *T*, and *a* and b are constants (Dunlop and Özdemir, 1997). This approach was necessary since the induced magnetization of paramagnetic constituents begins to saturate in high fields at low-temperatures and cannot be approximated with a linear function. Using this approach, excellent fits to the magnetization curves were achieved (Fig. 6).

The results of hysteresis measurements for the suboxic and anoxic samples are largely different. To represent the suboxic and



Fig. 7. Low-temperature evolution of hysteresis loops for magnetic extracts from samples of the upper suboxic (a and b) and lower anoxic (c and d) core sections. For the sake of clarity, hysteresis loops at six selected temperatures are shown, indicated by small numbers in the left side panel. Data shown on the left are uncorrected, data on the right are corrected for dia- and paramagnetic contributions.

Fig. 8. Results of normalized hysteresis parameters: low-temperature dependence of (a) saturation magnetization $M_{5,}$ (b) saturation remanent magnetization M_{RS} , and (c) coercive field B_{C} .



Та	bl	e	4	

m								P
lemperature-d	lependent va	alues for hyste	resis parameter	s saturation ma	agnetization M_{S} ,	saturation remanence M	ing and coercive for	$ce B_C$

T [K]	[K] 1.65 m		2.95 m			13.25 m	14.55 m		14.55 m	14.55 m			15.25 m		
	$M_{\rm S} [10^{-6} {\rm Am}^2]$	$M_{\rm RS} [10^{-6} {\rm Am^2}]$	<i>B</i> _C [mT]	$M_{\rm S} [10^{-6} {\rm Am^2}]$	$M_{\rm RS} [10^{-6} {\rm Am}^2]$	<i>B</i> _C [mT]	$M_{\rm S} [10^{-6} {\rm Am}^2]$	$M_{\rm RS} [10^{-6} {\rm Am}^2]$	<i>B</i> _C [mT]	$M_{\rm S} [10^{-6} {\rm Am}^2]$	$M_{\rm RS} [10^{-6} {\rm Am}^2]$	$B_{\rm C} [{\rm mT}]$	$M_{\rm S} [10^{-6} {\rm Am}^2]$	$M_{\rm RS} [10^{-6} {\rm Am^2}]$	<i>B</i> _C [mT]
295	3.98	0.67	12.56	1.59	0.25	10.31	0.78	0.13	9.47	0.36	0.05	6.80	0.46	0.06	9.17
245	3.98	0.74	15.05	1.71	0.30	12.96	0.93	0.19	13.56	0.42	0.07	10.15	0.52	0.09	12.87
195	4.02	0.81	17.54	1.72	0.35	16.28	1.06	0.26	19.65	0.48	0.10	14.82	0.58	0.12	18.97
145	4.00	0.87	19.98	1.74	0.39	20.67	1.19	0.34	29.77	0.51	0.12	21.79	0.65	0.17	28.32
110	4.02	0.91	22.19	1.76	0.43	24.89	1.28	0.40	40.72	0.53	0.15	29.44	0.69	0.20	38.54
75	4.04	0.98	25.54	1.75	0.47	30.74	1.35	0.48	56.32	0.56	0.18	39.62	0.73	0.24	52.20
60	4.03	1.01	27.05	1.75	0.49	33.58	1.37	0.52	65.08	0.57	0.20	45.40	0.74	0.26	60.00
50	4.02	1.03	28.30	1.75	0.50	35.96	1.39	0.55	73.26	0.57	0.21	50.55	0.75	0.28	68.20
40	4.03	1.08	30.56	1.75	0.53	39.82	1.41	0.58	84.67	0.58	0.22	57.60	0.76	0.29	77.89
35	4.04	1.12	31.93	1.75	0.55	41.95	1.43	0.61	93.12	0.59	0.23	62.39	0.76	0.30	84.98
30	4.05	1.18	34.41	1.76	0.57	46.09	1.45	0.63	104.25	0.59	0.24	69.05	0.77	0.32	93.69
25	4.10	1.23	36.93	1.78	0.60	49.68	1.47	0.67	118.51	0.60	0.25	78.13	0.78	0.33	104.76
20	4.12	1.27	39.73	1.80	0.61	54.41	1.49	0.70	135.97	0.60	0.27	87.98	0.79	0.35	118.60
15	4.19	1.29	43.60	1.84	0.62	57.71	1.51	0.74	163.73	0.62	0.28	102.12	0.81	0.37	140.50
10	4.36	1.28	46.65	1.89	0.63	62.33	1.58	0.80	195.91	0.64	0.30	119.85	0.83	0.39	171.30
5	4.60	1.30	49.81	1.98	0.65	66.60	1.68	0.86	233.50	0.67	0.31	140.45	0.89	0.42	205.10

anoxic environments, the temperature evolution of the hysteresis loops is shown for the samples from 1.65 m (Fig. 7a) and 15.25 m (Fig. 7b) depth. For the sake of clarity only six selected temper-

hysteresis loops systematically change in shape with decreasing ature steps (295, 195, 110, 50, 20 and 5 K) are given here. The

temperature and increasing sediment depth. At room-temperature

be compared with each other. The absolute values are summarized normalized to their respective value at 5 K so that the samples can to the inability of mass determination, the parameters have been tified after dia- and paramagnetic correction is shown in Fig. 8. dependence of the coercive force $B_{\rm C}$ (Table 4).

The temperature dependence of the hysteresis properties quan-

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temperatures. This effect is most pronounced in the temperature samples from the anoxic zone exhibit distinct hysteresis at lower the loops become more pronounced and wider, in particular, the a soft ferrimagnetic mineral phase. With decreasing temperature moderate fields (<500 mT), which is indicative of the presence of coercivities. Saturation magnetization is usually reached in low to hysteresis loops for all extracts are quite narrow and yield low The saturation magnetization M_S (Fig. 8a) for the suboxic samples shows only subtle temperature variations between 295 and 30 K. Upon further cooling to 5 K an increase of about 20% is observed. In contrast a quasi-continuous increase for M_S with decreasing temperature can be seen for the anoxic samples. At 5 K the values for M_S are about a factor of two to three higher for the anoxic and only 1.15 for the suboxic samples compared to room temperature. However, the complexity of the paramagnetic correction may effect the accurate determination of M_S and thus the results should be treated with caution.

Saturation remanence $M_{\rm RS}$ (Fig. 8b) for the suboxic samples increases slightly between 295 and 50 K and in a steeper manner below 50 K. $M_{\rm RS}$ seems to reach saturation at very low temperatures, however, the overall increase of two to three times is relatively small. For the anoxic samples a continuous increase with decreasing temperature is observed, with an overall 7-fold increase between 295 and 5 K.

At room temperature, values of B_C (Fig. 8c) range between 6 and 13 mT for all samples. Thompson and Oldfield (1986) report this as a typical value for multi-domain magnetite. Upon cooling, B_C varies little between 295 and 50 K. Below 50 K the coercive field starts to increase sharply, reaching values of 50–67 mT at 5 K for the suboxic samples and up to 140–233 mT for the anoxic samples. This equals a 4- and 7-fold increase for the suboxic samples and on average 25-fold higher coercivities for all of the anoxic samples. No ordering temperature could be determined from the temperature dependence of the coercivity (such as the magnetic ordering transition from ferrimagnetism to paramagnetism at 210 K for TH80, as seen before in the RT-SIRM cycling experiments, compare Fig. 5). The distinct increase of B_C below 50 K might potentially be due to the hemoilmenite entering its spin glass state (Burton et al., 2008).

The hysteresis loops at certain low-temperatures exhibit a peculiar wasp-waisted shape, which is more pronounced for the anoxic than for the suboxic samples (Fig. 9). Roberts et al. (1995) and Tauxe et al. (1996) describe this phenomenon as indicative for a mixture of relatively high- and low-coercive material. Such a mixture could consist of either distinct magnetic minerals or a bi-modal grain-size distribution of a single magnetic component.

Large quantities of superparamagnetic magnetite or hard coercivity particles (such as hematite or goethite) are commonly assumed to be the main reason for wasp-waisted hysteresis loops. In this study these options appear implausible for several reasons: the applied extraction method tends to favor the collection of rather low-coercivity components (Franke et al., 2007b). Moreover, hematite and/or geothite must be a hundred times more abundant in the sediment to rival the magnetization of (titano-)magnetite. Secondly, the observed hysteresis loops become more constricted with decreasing temperature. Superparamagnetic (magnetite) particles are expected to cause constriction in the room-temperature loops rather than low-temperature loops because they block at lower temperatures and become stable single domain grains (Dunlop, 1973).

An alternative explanation for the wasp-waisted hysteresis loops at low-temperature are the observed Ti-rich hemoilmenites with compositions 0.85 < y < 1. These members of the ilmenite– hematite solid solution series are paramagnetic at room temperature but exhibit superparamagnetic properties just below room temperature and their Néel temperature (Ishikawa, 1962) and antiferromagnetic behavior at lower temperatures (Burton et al., 2008).

5. Discussion

SEM analyses have shown, that the predominant magnetic minerals in the upper suboxic sediments are (titano-)magnetites of variable Ti composition. Ilmenohematites and hemoilmenites are also present, but relative to (titano-)magnetite they are less abundant and thus less important in this depth interval. Dillon and Bleil (2006) already suggested that the vast majority of the magnetic assemblage reaching the gravity core location was composed of eroded volcanic material originating from the Cameroon Volcanic Line. Herrero-Bervera et al. (2004) and Ubangoh et al. (2005) reported titanomagnetites, both poor and rich in Ti, as the principal magnetic minerals in rocks from the Cameroon volcanics in the drainage area of the Niger River tributaries. Ubangoh et al. (2005) also observed rocks with abundant ilmenite lamellae in the titanomagnetites. Both investigations used Curie temperature experiments above room temperature for the discrimination of the magnetic particles. Hemoilmenites with high Ti-contents could not be determined by these former studies, as they may actually have Curie temperatures below room temperature. Combined with our microscopy findings, which show that hemoilmenites are a minor component of the primary magnetic mineral assemblage compared to (titano-)magnetite, sub-room-temperature Curie points provide a possible explanation why members of the ilmenite-hematite solid solution series have not been identified previously.

In the lower anoxic sediments early diagenetic alteration after deposition has resulted in a more complete depletion of the (titano-)magnetite phase, producing a relative enrichment of the hemoilmenite phase in the strongly diagenetically altered section so that the relative abundance of the hemoilmenite particles increases to \sim 66%. This change in abundance can be interpreted in terms of selective diagenetic effects on the magnetic mineral assemblage. Dillon and Bleil (2006) reported that reductive diagenesis has extensively affected the magnetic mineral inventory of these organic-rich late Quaternary sediments in the Niger deep-sea fan. Fig. 2b shows the extent of the suboxic zone and the transition to anoxic conditions in the pore water profile of the sedimentary sequence. The rock magnetic investigation showed that the diagenetic processes did not act uniformly across the detrital magnetic mineral assemblage, but instead depended strongly on particle size and Ti-content, with the smallest particles dissolving first and the Ti-rich component being the most resistant to the reducing conditions. Another important determining factor is the availability of sulfur in this geochemical system.

Due to the already considerable strong alteration of the upper suboxic sediments, one of the main challenges in this study is the generally low magnetic mineral content. Therefore the concentration of magnetic carriers is even lower in the anoxic sediments and only relicts of the primary magnetic mineral assemblage remain. Highly sensitive low-temperature magnetic measurements on the magnetic extracts were therefore required for magnetic mineral identification. These measurements revealed large contrasts between the samples from the suboxic and anoxic layers. This emphasizes that the Fe–Ti oxide phases occur in different proportions in the two different geochemical zonations, which is also reflected in the magnetic behavior of these samples.

5.1. Suboxic samples

In the upper suboxic sediments (titano-)magnetite particles dominate the low-temperature magnetic characteristics. The increase in the RT-SIRM cooling curves just below roomtemperature and the broad decrease around 120 K were described by Özdemir and Dunlop (2003) and Kosterov (2003) as characteristic behavior for a mineral assemblages dominated by (Ti-poor) titanomagnetite (Fig. 5). Additionally, the low-temperature hysteresis measurements confirm the presence of a soft ferrimagnetic mineral phase, documented in the relatively small increase in hysteresis parameters with decreasing temperature. The value of the coercive force, showing an overall increase by a factor of four to



Fig. 9. Uncorrected hysteresis loops measured at 50 K; suboxic samples (a) and (b) show slightly wasp-waisted characteristics, whereas anoxic samples (c) to (e) show a distinct wasp-waisted shape.

seven between 300 and 5 K, is comparable with results obtained from synthetic magnetites by Özdemir et al. (2002). Harrison and Putnis (1995) analyzed the low-temperature dependence of the magnetite-ulvöspinel solid solution series and observed low coercivities (10 mT) for the magnetite-rich (Fe-rich) end-members at low-temperatures (4.4 K). The synthetic submicron magnetites in the study of Özdemir et al. (2002) yield coercivities of \sim 15 mT at room temperature and \sim 50 mT at 15 K, which is a similar increase in $B_{\rm C}$ by a factor of three to four. The coercivity of the synthetic samples increases sharply when crossing the Verwey transition because their compositions are close to stoichiometry. The lack of the Verwey transition in the suboxic samples of this study is presumably due to their Ti-content, since at fairly low Ti-contents of x > 0.04 the transition is suppressed (Kakol et al., 1994). The small amount of observed shrinking cracks is indicative of a small degree of maghemitization, which may additionally subdue the Verwey

transition (Dunlop and Özdemir, 1997). Tucker (1981) reported on low-temperature magnetic hysteresis properties for two multi domain titanomagnetite crystals with compositions of x = 0.52 and x = 0.65. The first titanomagnetite (x = 0.52) shows values of $B_{\rm C}$ $(\sim 25 \text{ mT})$ and $M_{\text{RS}}/M_{\text{S}}$ (0.29) at 77 K that are comparable with our results at 75 K of $B_{\rm C}$ = 25 mT and a $M_{\rm RS}/M_{\rm S}$ ratio of 0.25. The calculated average titanomagnetite composition for our suboxic mineral assemblage results in a mean value of TM55 which is very close to the composition of the respective single crystal of Tucker (1981). For the TM65 crystal from the study of Tucker (1981) the reported values are higher ($B_{\rm C}$ = 58 mT; $M_{\rm RS}/M_{\rm S}$ = 0.58), with the increased coercivity explained as a result of increasing in magnetic anisotropy. At even lower temperatures of 4.2 K, Schmidbauer and Readman (1982) observed coercivity values of comparable magnitude for titanomagnetites with compositions of x = 0.5 ($B_C \sim 31$ mT) and $x = 0.6 (B_{\rm C} \sim 70 \,{\rm mT})$.



Fig. 10. Low-temperature hysteresis loops for synthetic sample 6F72x2.4 measured at 55 K (black solid line) and 5 K (grey solid line), showing slightly (55 K) and pronounced (5 K) wasp-waisted shapes (Engelmann et al., unpublished data).

The hysteresis loops of the suboxic samples show wasp-waisted shapes at lower temperatures. The presence of an intermediate titanomagnetite phase in paragenesis with a hemoilmenite phase as described by Krása et al. (2005) would offer a reasonable explanation for this behavior. According to Buddington and Lindsley (1964) the ilmenite content of a hemoilmenite phase coexisting with a titanomagnetite phase with a composition of x > 0.5 would be y > 0.85. Since hemoilmenite phases of y > 0.87 exhibit superparamagnetic properties at low-temperatures (Ishikawa, 1962), this could explain the constriction that was observed in the hysteresis loops of the suboxic samples. This is also consistent with the SEM and EDS analyses, which revealed the presence of hemoilmenite as minor constituents ($\sim 17\%$) in the primary magnetic mineral assemblage of the suboxic samples.

Nevertheless, the Ti-contents derived from the SEM analysis are not overly high. However, the hysteresis loop for the synthetic sample 6F72x2.4 clearly shows wasp-waisted characteristics at temperatures \leq 55 K (Fig. 10), even though the composition of the hemoilmenite phase is below *y* = 0.8. With *y* = 0.76 the hemoilmenite composition of 6F72x2.4 is still higher than in our natural sample, but it generally shows that wasp-waisted shapes do occur for such Ti-contents. The distribution of the two phases in the synthetic sample is comparable (79% TM: 21% IlmHem) to the derived distribution in our natural suboxic sample (82% Mt/TM: 17% IlmHem). Therefore the wasp-waisted shape in our hysteresis loops is most likely the result from the interplay between titanomagnetite and hemoilmenite.

5.2. Anoxic samples

The results of the low-temperature measurements from the anoxic layers lead to the conclusion, that titanomagnetite cannot be the predominant magnetic mineral phase in this depth interval. Here, Ti-rich hemoilmenite dominates the magnetic mineral assemblage. This was also confirmed by the findings of Heslop and Dillon (2007), who used a non-negative matrix factorization (NMF) algorithm to unmix remanence data into constituent end-members from the same Niger deep-sea fan sediments. Based on the coercivity distribution, fine and coarse grained (titano-)magnetite was found to make the greatest contribution to the remanence signal in the upper sediments. However, a third component was needed to successfully model the detrital magnetic mineral assemblage. This third component comprised the hardest coercivity and was argued to be Ti-rich titanomagnetite or hemoilmenite. It could be shown by the present study, that below the anoxic/sulfidic transition, this Ti-rich component had the highest resistance to reductive dissolution and dominates the magnetic mineral assemblage. The coarse and fine (titano-)magnetite fraction almost disappears completely towards the base of the sediment core.

During cooling, the RT-SIRM increases continuously with decreasing temperature. Franke et al. (2007c) considered two possibilities for this phenomenon, either the presence of a high-coercive mineral such as goethite or a Ti-rich oxide phase. Due to their assumption that high-coercivity minerals are generally underrepresented in magnetic extracts, it appears more probable for the natural samples in this study that a Ti-rich mineral phase is responsible for the observed increase of RT-SIRM during cooling. The assumption that this behavior can be attributed to the presence of hemoilmenite in our Niger deep-sea fan samples is therefore reasonable, although the interpretation of low-temperature RT-SIRM curves from comparable natural samples is still lacking in the literature (Kosterov, 2007).

Fig. 11 compares the low-temperature measurements performed on the natural sample from gravity core GeoB 4901 (15.25 m; Fig. 11a) and the well-defined synthetic sample 6F72x2.4 (Fig. 11b). The continuous increasing trend in the RT-SIRM curves with decreasing temperature can be seen for both samples. Since the composition of the synthetic sample is well known, goethite can be excluded as a possible reason for this low-temperature behavior. Here this phenomenon can be rather attributed to the presence of a Ti-rich mineral phase. The RT-SIRM curve of another welldefined single phase titanomagnetite of synthetic origin (TM60) (not shown here; Engelmann, unpublished data) shows a rather similar RT-SIRM curve progression as the suboxic samples of this study, with a large decrease around 190 K. This particular single phase sample does not contain any hemoilmenite. Therefore we suggest that in the two-phase synthetic sample 6F72x2.4 the distinct low-temperature increase refers to the presence of the Ti-rich hemoilmenite phase.

This sample possibly shows paramagnetic-like LT-behavior caused by the presumable presence of near end-member ilmenite. The magnetically harder nature of this mineral phase suggests that a higher coercivity magnetic component other than (titano-) magnetite must be present to create such a low-temperature increase in the RT-SIRM. In particular the EDS results of the SEM analyses confirm an overall higher content of Ti-rich particles in the lower anoxic part of the sediment series. The main component here is hemoilmenite (\sim 66%) with compositions between Ilm50Hem50 up to nearly pure ilmenite (Ilm99Hem01).

The unusually high coercivities of the anoxic samples at lowtemperatures also concur with the presence of hemoilmenite, rather than titanomagnetite. The stability at low-temperatures with $B_{\rm C}$ values up to 220 mT is remarkable. For titanomagnetites such high coercivities at low-temperatures have been rarely published in the literature. Schmidbauer and Readman (1982) reported coercive forces of 500 mT for synthetic titanomagnetites with a composition of x = 0.8 at 4.2 K, which is twice as high as the natural samples in our study.

Brown et al. (1993) also reported coercivity values for members of the ilmenite-hematite solid solution series. For synthetic ilmenohematites (=titanohematites) with compositions of y=0.2and y=0.4 to 0.6 they observed values between 170 and 260 mT at temperatures of 125 K. They also found that samples quenched at high-temperatures have significantly higher coercivities than those annealed at temperatures below the order-disorder transition of hemoilmenite. Nord and Lawson (1992) correlate this behavior to the development of twin domain boundaries during quenching from high-temperature, and suggest that annealing above the order-disorder transition temperature results in a substantial increase in twin-domain boundaries and a drastic increase in the coercivity of the material.



Fig. 11. Low-temperature variation of RT-SIRM for (a) the magnetic extract from 15.25 m depth and (b) the synthetic sample 6F72x2.4 (Engelmann et al., unpublished data).

Gehring et al. (2007) studied micromagnetic properties of ilmenite–hematite particles originating from an alluvial soil. X-ray diffraction revealed hemoilmenite grains with an ilmenite mole fraction of y = 0.86. The hysteresis loops at 20 K yielded $B_C = 92$ mT, reached $B_C = 144$ mT at 15 K, and $B_C = 244$ mT at 6 K. These values are clearly consistent with the results of our study (compare Table 2). Here, the calculated average hemoilmenite composition is Ilm77Hem23. In metamorphic ilmenite–hematite bearing rocks from Norway stable natural remanences have been identified by McEnroe et al. (2001). Such natural phases show high coercivities that are attributed to their small grain-sizes (Merrill, 1968; McEnroe et al., 2001).

Besides the unusually high coercivities of the natural samples, the wasp-waisted shape of the low-temperature hysteresis loops hints at the presence of Ti-rich hemoilmenite phases. For the suboxic samples the wasp-waisted shape could be explained by the interaction between titanomagnetite and hemoilmenite. In the anoxic sediments titanomagnetite is only present in very minor amounts (\sim 5%), but wasp-waisted hysteresis loops are still observed. These are most pronounced in the temperature interval between 30 and 50 K. Apparently, another mechanism must be responsible for this phenomenon. The hysteresis loops measured by Gehring et al. (2007) showed typical wasp-waisted shapes at temperatures between 150 and 30 K, which became more pronounced with decreasing temperature. They explained the wasp-waisted shape by the generation of short-range ordered areas in the super-paramagnetic state within the hemoilmenite particles.

Hemoilmenites with $0.8 \le y \le 0.87$ are known to exhibit superparamagnetic behavior at temperatures below 50 K (Ishikawa et al., 1985) and thus can account for the wasp-waisted shape of the hysteresis loops at low-temperatures. Such Ti-rich phases are rather untypical as individual homogenous grains, but are more likely to be present as intergrown exsolution lamellae within the Ti-poor ilmenohematite matrix. Dunlop and Özdemir (1997) report that if the cooling of a melt is slow enough, hemoilmenite of intermediate composition tends to exsolve into intergrown Ti-poor (near hematite) and Ti-rich (near ilmenite) phases. However, compositions with y > 0.8 were identified in the magnetic extracts of this study. Whether they consist of either homogenous grains or of intergrown exsolved phases cannot be verified by the technique used for our unconsolidated samples but it seems very likely.

Nord and Lawson (1992) reported that hemoilmenites with y > 0.5 are magnetically inhomogeneous and consist of cationordered ferrimagnetic microstructures which are separated by boundary layers. Earlier, Lawson and Nord (1984) showed that synthetic Ilm80Hem20 (y = 0.8) samples had remanence properties that can be explained by the presence of a single SD-like magnetic carrier, even though Ilm80Hem20 should be paramagnetic at room temperature. The source of the measured remanence is argued to be SD-like material within the grains themselves, resulting from the generation of transformation induced domain boundaries, which are partly enriched in the hematite component relative to the bulk composition of the grain.

Specific low-temperature magnetic measurements can be used to differentiate between Fe and Ti dominated magnetic mineral assemblages. If Fe-rich mineral phases prevail, RT-SIRM cycles in the low-temperature range show a distinct decrease around the Néel temperature and/or Verwey transition of magnetite. For Ti-rich particles the RT-SIRM curves continuously increase with decreasing temperature. However, solely on the basis of the RT-SIRM experiments it is not possible to differentiate unambiguously between the magnetite-ulvöspinel ($Fe_{3-x}Ti_xO_4$) and ilmenite-hematite solid solution series (Fe_{2-y}Ti_yO₃). The hysteresis parameters reveal additional diagnostic features that can help to differentiate between Fe-rich and Ti-rich oxide mineral phases. The characteristic evolution of the measured loops with decreasing temperature points towards two distinct mineral phases in the different environments. In particular the coercive force was found to be a helpful indicative parameter, with a 25-fold increase characteristic of phases from the ilmenite-hematite solid solution series. Such high values were not found for any Fe-rich mineral phase. The high coercivities most likely result from exsolution or lamellar magnetism in hemoilmenite grains.

5.3. Environmental implication: correlation of Fe and Ti vs. κ

EDS analyses have clearly shown, that Fe-rich particles dominate the magnetic mineral assemblage in the upper suboxic environment, whereas Ti-rich particles prevail in the lower anoxic section. The dominance of Ti-rich magnetic particles in the lower anoxic environments is also strongly reflected in the downcore profile of the volume magnetic susceptibility κ . Fig. 12 (left) shows the downcore profiles of κ compared to the solid phase concentrations of Fe and Ti in the bulk sediments derived from ICP-EAS analysis (Zabel et al., 2001).

The correlation between κ and Fe is in general clear throughout the whole depth profile, as is the correlation between κ and Ti. Only for the depth interval from the top of the core to the modern Fe-redox boundary (upper three meters) the correlation is not significant. However, for this section the correlation is higher between κ and Fe (r=0.17, Pearson; Fig. 12d) relative to the correlation between κ and Ti (r=0.05; Fig. 12e). Below the modern Fe-redox boundary the correlation between κ and Ti is consistently higher than between κ and Fe. Not including the upper three meters, Pearson's correlation coefficient between κ and Ti (r=0.70) is noticeably higher than between κ and Fe (r=0.51).



Fig. 12. (Left) downcore profiles of (a) volume magnetic susceptibility κ and solid phase concentrations of (b) Fe and (c) Ti (Zabel et al., 2001). All parameters are plotted on a linear depth scale and a non-linear age scale (Adegbie, 2001) including marine oxygen isotope stages (MIS) and terminations (TI and TII). Grey shading indicates cold periods. For further details see also Fig. 2. (Right) correlation of (d) κ vs. Fe-content, and (e) κ vs. Ti-content. Symbols indicate data points for the upper 3 m (black triangles), the depth interval from 3.0 m to 12.5 m (open grey diamonds) and below 12.5 m (filled black diamonds). Pearson's correlation coefficients *r* are given for the respective depth intervals.

Between 3.0 and 12.5 m core depth the correlation of κ and Ti is considerably higher (r=0.79) than for κ and Fe (r=0.29; Fig. 12d and e). This suggests that below the modern Fe-redox boundary the relative contribution of Ti-rich mineral phases to the magnetic susceptibility signal increases notably. The transition from suboxic to anoxic sedimentary conditions is located at 12.5 m core depth. Below this transition the highest correlation of r = 0.86 exists between κ and Ti. For the same depth interval the correlation also increases slightly again for κ and Fe (r=0.59; Fig. 12d). Here, iron is mainly bound to non-magnetic phases such as pyrite, which was identified in previous high-temperature thermomagnetic measurements (Dillon and Bleil, 2006). This agrees well with the fact that the magnetic susceptibility of hemoilmenites $(\chi = 100 - 100000 \times 10^{-8} \text{ m}^3/\text{kg}; \text{Bleil and Petersen}, 1982)$ is at least one magnitude higher compared to the magnetic susceptibility of pyrite ($\chi = 30 \times 10^{-8} \text{ m}^3/\text{kg}$; Thompson and Oldfield, 1986).

In the uppermost core section above the modern Fe-redox boundary, the good correlation between κ and Fe implies that reactive Fe is bound in Fe-rich magnetic mineral phases. These phases dominate the magnetic mineral assemblage, both in concentration and concerning their magnetic properties.

In the subsequent lower suboxic and anoxic intervals the clear correlation between κ and Ti and the less pronounced correlation between κ and Fe demonstrates that most of the Fe is bound to phases that are non-magnetic at room temperature or is present in its reduced state. Here reactive Fe was subsequently depleted, relocated and the formation of secondary Fe minerals took place. As a consequence the influence of the Ti-rich phases on the magnetic signal increases significantly. This is also reflected in the magnetic susceptibility record. Therefore we conclude that the degree of correlation between κ and Fe, and κ and Ti, represents whether the magnetic assemblage is dominated rather by Fe-rich magnetic min-

eral phases, components of intermediate Ti-content, depleted in reactive Fe, or by Ti-rich magnetic phases, including mainly nonreactive Fe.

6. Conclusions

In this study the different magnetic components in sediments from the Niger deep-sea fan were determined in detail by a combination of low-temperature measurements with scanning electron microscopy analyses applied to magnetic extracts from various geochemical zones.

The magnetic analyses indicate a fairly complex mineralogy of the primary magnetic mineral assemblage, which is dominated by (titano-)magnetite of variable composition derived from the Cameroon volcanics in the drainage area of the Niger River tributaries (Herrero-Bervera et al., 2004; Ubangoh et al., 2005). Above the modern redox boundary hemoilmenite originating from the same source area is also present. Magnetically this phase does not play such an important role here because the concentration is too low and the magnetic moment is too weak.

Nevertheless, below 3.0 m core depth, processes of reductive diagenesis start to influence the composition of the magnetic mineral assemblage. The portion of Fe-oxides decreases almost to zero and Ti-bearing mineral phases become more important in this depth interval. Although hemoilmenites are abundant, titanomagnetites are still the major contributor to the magnetic signal.

Below the sulfidic transition, located in a depth of 12.5 m, grains of the ilmenite–hematite solid solution series are found to be the prevailing contributor to the magnetization of the anoxic sediments. In the course of reductive diagenesis, alteration has resulted here in a more complete depletion of titanomagnetite. (Titano-) magnetite is most likely replaced by paramagnetic iron sulfide minerals. Hemoilmenite is therefore much more resistant to reductive diagenesis due to its low content of reactive iron.

The study has also shown that even when remanence transitions and susceptibility peaks are not obviously apparent in low-temperature data – which is the case for many marine sediment samples – careful investigations of the data reveal important relevant information. This can be used to characterize magnetic mineral phases in a sediment sample and results therefore in a detailed paleoenvironmental understanding.

Acknowledgements

The synthetic sample 6F72x2.4 and the respective unpublished data were provided by Ralf Engelmann and Dominque Lattard (University Heidelberg, Germany), the authors are grateful for their support and various discussions. We would like to thank T. Frederichs (University of Bremen, Germany) for technical assistance and annotation of the low-temperature MPMS analyses. The SEM and EDS analyses were performed at the EMU (Utrecht University, the Netherlands). The authors would like to thank F. Lagroix and an anonymous reviewer for their helpful comments and U. Bleil for constructive criticism of the manuscript. This study was funded by the DFG.

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