EPR evidence for maghemitization of magnetite in a tropical soil

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SUMMARY
Electron paramagnetic spectroscopy (EPR) was used in combination with standard rock magnetic methods to study magnetic minerals in a tropical soil. The susceptibility and hysteresis measurements showed magnetite grains with a Curie temperature near 850 K as the dominant magnetic remanence carriers in the soil. A minor Ti content in the magnetite was found by energy dispersive X-ray analysis. In order to get insight into the weathering status of the magnetite, different chemical treatments, including oxalate and citrate–bicarbonate–dithionite (CBD) extraction, were applied to the soil samples. The hysteretic properties exhibited no significant differences between the untreated and the CBD or oxalate treated samples. By contrast, the comparison of the EPR spectra revealed a significant broadening of the linewidth (δB) and a shift of the g-values (g_{eff}) to lower fields after the CBD treatment. Furthermore, the spectral parameters g_{eff} and δB exhibited an angular dependence. At low temperature, the CBD treated samples showed a jump in δB between 120 and 100 K, the temperature range characteristic for the Verwey transition in magnetite. The changes in the spectral properties after the CBD treatment, which dissolves ferric oxides, were attributed to the removal of maghemite formed by the oxidation of magnetite, that is, during the maghemitization of the magnetite grains.

Key words: EPR spectroscopy, FMR, magnetite, maghemitization, weathering.

1 INTRODUCTION
Lithogenic magnetite is widespread in soils and often dominates their magnetic properties. During weathering, magnetite becomes oxidized, whereas under reducing conditions, as in waterlogged soils, it can undergo reductive dissolution (e.g. Henshaw & Merrill 1980). The identification of such processes is important to deduce pedogenesis. Oxidation of magnetite can lead to the formation of maghemite or less crystallized ferric phases. Using XRD and microscopic analysis, Gallagher et al. (1968) could show on a well defined synthetic magnetite sample a topotactical conversion into maghemite. The conversion is known as maghemitization, and has been described also in natural systems such as pelagic sediments (Smirnov & Tarduno 2000) and loess (Chen et al. 2005). In the first study, maghemite was detected by thermal demagnetization, while in the second one it was observed under the transmission electron microscope. Under reducing conditions, magnetite is slowly dissolved. The rate is grain size dependant and is generally much slower than for ferric oxides.

The aforementioned redox processes are surface controlled (Stumm & Morgan 1996). The detection of the surface products with rock magnetic methods is difficult, because the magnetic signals are generally dominated by the bulk properties. Furthermore, magnetite and its oxidation product maghemite have similar magnetic properties (Dunlop & Özdemir 1997). In the rock magnetic literature, there are several experimental approaches to detect maghemitization. Van Velzen & Dekkers (1999) proposed that heating to 150°C could be used to distinguish between maghemite and partially oxidized magnetite because such a treatment would selectively change the magnetic coercivity of the latter. Low temperature magnetic experiments offer another approach, in which the oxidation of magnetite is detected via the suppression of the so-called Verwey transition (Özdemir et al. 1993). This transition, however, is also suppressed in magnetite with Ti substitution (Kakol et al. 1992); therefore, ambiguity can remain when using such an approach.

Electron paramagnetic resonance (EPR) spectroscopy is a powerful tool for studying paramagnetic cations such as Fe^{III} in diluted systems (e.g. Meads & Malden 1975; Angel & Vincent 1978). A detailed description of the technique is given by Wertz & Bolton (1972). The method, however, has rarely been used to study phases carrying a remanent magnetization. In such systems, strong dipole–dipole interactions or magnetic coupling can lead to intense and broad signals at room temperature (RT) that cannot be assigned unambiguously to a specific magnetic phase (Griscim 1980; Gehring & Karthein 1990). Such broad signals are produced by ferromagnetic resonance (FMR). The basic principles of this phenomenon were described by several authors (Kittel 1948; Bickford 1950). Experimental studies on magnetite revealed that apart from the different anisotropies (e.g. magnetocrystalline) the skin effect can influence the FMR (Kittel 1946). Such an effect is produced in electrical conductors where eddy currents limit the penetration depth of the electromagnetic radiation. For magnetite at RT, this depth is...
in the \( \mu \text{m} \) range, and, therefore, is expected for multidomain (MD) and larger pseudo-single domain (PSD) particles. By contrast, in single-domain (SD) particles, the electromagnetic radiation is uniform across the whole grain. Hence, for MD and larger PSD grains, the surface and the near-surface region contribute more strongly to FMR spectra.

With this in mind, it is conceivable that FMR properties of larger magnetite grains can be influenced by surface modifications caused by oxidation. In order to test this hypothesis, we applied EPR spectroscopy in concert with rock magnetic methods to lithogenic magnetite in a tropical soil which was subjected to different chemical treatments to selectively remove certain minerals or surface layers formed during weathering.

2 METHODOLOGY

2.1 Sample preparation

The soil samples were collected from the top layer of an alluvial Vertisol in the Savanna woodland of southern Mali near the hamlet Kabogora (Gehring et al. 1997). The lithogenic magnetite is dispersed in the soil matrix which mainly consists of quartz and montmorillonite. For the chemical and magnetic analyses, the samples were dried at 60 °C and passed through a 2-mm sieve in order to remove larger grains and components such as plant residues.

Three different chemical extraction procedures were performed on the untreated sample (KA). Oxalate treatment following the recipe by Schwertmann (1964) was used to remove amorphous and poorly crystallized ferric oxides from the soil samples. The citrate–bicarbonate–dithionite (CBD) extraction was performed using a protocol modified from Mehra & Jackson (1960). Briefly, 2 g of sieved soil, 2 g of sodium dithionite, 10 ml of 1 M sodium hydroxide solution, and 150 ml 0.3 M sodium citrate were equilibrated at RT for 24 h in dark bottles using an end-over-end shaker. The liquid phase was removed by filtration and the residue washed several times with doubly deionized water. The acid treatment involved boiling the sample in a concentrated acid solution with a 19:1 ratio of HCl to H\(_2\)SO\(_4\) followed by magnetic separation (Goulart et al. 1994). The treated samples are referred to as KA\textsubscript{ox}, KA\textsubscript{dith} and KA\textsubscript{acid}, respectively.

2.2 Experimental methods

The morphology of magnetic separates was investigated under the scanning electron microscope (SEM), which for semi-quantitative chemical analyses was connected to an energy dispersive system (EDX). The magnetic susceptibilities (\( \chi \)) of the untreated and chemically treated specimens were determined on an AGICO KLY-2 Kaplabor. In order to determine inhomogeneity effects, the susceptibility was measured on ten specimens. For sample KA, the temperature dependence of \( \chi \) between RT and 973 K was recorded with a heating rate of approximately 11 °C min\(^{-1}\). The hysteretic properties of all samples were analysed at RT by magnetization versus applied field experiments using a Vibrating Sample Magnetometer (VSM) from Princeton Corporation. Coercivity force (\( B_C \)), remanent magnetization (\( M_r \)) and saturation magnetization (\( M_s \)) were determined from the hysteresis loops, after they were corrected for paramagnetic contribution with a factor of 0.7. In addition, first-order reversal curves (FORC) were measured with the same instrument, and the data were analysed with a Matlab code by Winkelhofer (personal communication 2006).

The EPR spectra of the untreated and chemically treated samples (about 300 mg) were obtained at RT on a Bruker EMX 300 spectrometer. The X-band spectra were recorded at a microwave frequency of 9.77 GHz with a power of 0.2 mW, a modulation frequency of 100 kHz, and an amplitude modulation of 0.1 mT. The field strength was controlled by a Bruker NMR Gaussmeter ER 035M. In order to avoid movement of the magnetic particles due to alignment with the external field, the bulk sample in the glass tube was moulded in paraffin. The angular dependence of the FMR spectra was determined by rotating the glass tube in the EPR cavity with the help of a goniometer installed at the top of the cavity. The axis of rotation was perpendicular to the applied magnetic field. For each sample, the angular dependence was measured four times. The low temperature measurements were carried out on a Bruker Eleys spectrometer at a frequency of 9.5 GHz in a temperature range between 10 and 150 K. Prior to inserting the tubes into the spectrometer cavity, they were cooled in liquid nitrogen. The experimental parameters were the same as for the RT measurements.

2.3 FMR

Since FMR experiments have not often been used in geophysical research, the basics are summarized in the following paragraph. A detailed description can be found elsewhere (Kittel 1948; Vonsovskii 1966). In the presence of a magnetic field, the magnetic moments precess around the field direction with a frequency of \( \omega \). The frequency increases with the field, according to Larmor’s law:

\[
\dot{\omega} = \gamma B
\]

with \( \gamma \) being the gyromagnetic ratio, and \( B \) the applied field. Absorption occurs when this frequency is equal to the one of the microwave radiation (\( v \)). The resulting resonance condition can be written as

\[
hv = g\mu_B B
\]

assuming no anisotropy fields (\( h \) is Planck’s constant, \( g \) the splitting factor, \( \mu_B \) Bohr’s magneton). Since anisotropies in FM phases create internal fields which add to the external one, the effective field (\( B_{\text{eff}} \)) is the sum of applied field and internal fields. The \( g \)-value calculated from eq. (2) will, therefore, be modified to \( g_{\text{eff}} \). Eq. (2) becomes then

\[
hv = g_{\text{eff}}\mu_B B_{\text{eff}}
\]

Griscom (1980) describes four major anisotropy energies which can influence \( B_{\text{eff}} \). The magnetocrystalline anisotropy results from the crystallographic structure, whereas the shape anisotropy is caused by the interaction of the magnetization vector with the demagnetizing field of the grain. For magnetite, shape anisotropy is dominant in small (<20 \( \mu \text{m} \)) non-spherical particles, whereas for larger particles, magnetocrystalline anisotropy is prevailing (Dunlop & Özdemir 1997). Furthermore, domain structure anisotropy considers the relative orientation of the microwave field to the domain walls. Finally, exchange anisotropy arises from the interaction between magnetic particles.

For large magnetite grains at RT, the microwave radiation penetrates generally to a certain depth only, because of its relatively high electrical conductivity. The depth at which the intensity of the microwave radiation is reduced by 37 per cent is denoted the skin depth. The skin depth is frequency-dependent and it is about 5 \( \mu \text{m} \) for pure magnetite at the X-band frequency (Griscom 1974). Crystal defects or cation substitutions (e.g. Ti\(^{4+}\) for Fe\(^{3+}\)) often found in natural magnetites, however, can significantly reduce the electrical conductivity.
conductivity, and, therefore, increase the skin depth. Hence, spectroscopic information stems from both near-surface regions and the bulk of the larger magnetite grains.

The absorption spectra are generally presented as their first derivative. For the description of the FMR spectra of magnetite, two empirical parameters have been put forward: $g_{\text{eff}}$ value ($g$-value at zero-crossing), and the peak to peak linewidth $\delta B$. For non-interacting SD grains, these parameters can be used to characterize the anisotropy field. For systems with MD and PSD particles, however, these parameters can only be understood in a general sense.

3 RESULTS
3.1 Magnetic and microscopic properties
The mean susceptibility of KA was $69 \times 10^{-8}$ m$^3$ kg$^{-1}$ with a standard deviation ($\sigma$) of $4 \times 10^{-8}$ m$^3$ kg$^{-1}$. A mean $\chi$ of 77 ($\sigma = 6) \times 10^{-8}$ m$^3$ kg$^{-1}$ was found for KA$_{\text{ox}}$, while that of KA$_{\text{dith}}$ was 54 ($\sigma = 2) \times 10^{-8}$ m$^3$ kg$^{-1}$. Upon heating KA to 973 K, $\chi$ remained relatively constant up to 510 K followed by a bump at 575 K and a decrease up to 775 K. A more pronounced decay was found between 793 and 850 K (Fig. 1). At higher temperature $\chi$ did not significantly change any more. Upon reversed cooling, $\chi$ increased drastically down to about 520 K followed by a slight decrease. The end value at RT was three times higher than the starting value.

The magnetization versus applied field plot (Fig. 2) revealed an open hysteresis loop for KA, KA$_{\text{ox}}$ and KA$_{\text{dith}}$. By contrast, no hysteretic properties were found for KA$_{\text{acid}}$ (Fig. 2). The coercivity field ($B_c$) was 10 mT for KA and KA$_{\text{dith}}$, and 9 mT for KA$_{\text{ox}}$. The hysteresis loops for KA, KA$_{\text{ox}}$ and KA$_{\text{dith}}$ closed between 250 and 300 mT. The ratio of the remanent magnetization ($M_r$) to the saturation magnetization ($M_s$) was 0.15 for KA, 0.18 for KA$_{\text{ox}}$, and 0.17 for KA$_{\text{dith}}$.

The FORC diagram of KA revealed two features. The distribution along the $B_u$ axis extended up to 50 mT with weakly diverging open contours (Fig. 3a); a maximum was observed at $B_u \approx 10$ mT. The second feature consisted of a distribution along the $B_u$ axis for values of $B_u < 5$ mT. Chemical treatment did not significantly change the diagrams and both KA$_{\text{ox}}$ and KA$_{\text{dith}}$ displayed the same features as KA (Figs 3b and c).

Under the SEM, the magnetite grains were generally larger than 20 $\mu$m and exhibited variable morphologies. No significant change in the morphology was observed after chemical treatment (Fig. 4).

3.2 EPR measurements
The EPR spectra of the untreated, oxalate and the CBD treated samples recorded at RT were dominated by a broad signal (Fig. 5). This signal was missing in the KA$_{\text{acid}}$ sample. All samples exhibited two additional signals, one at low field with $g = 4.3$ and the other at high field with $g = 2$. The resonance at low field could be assigned to Fe$^{3+}$ in distorted octahedral coordination sites, whereas the sharp signal at $g = 2$ is characteristic for a free radical (e.g. Meads & Malden 1975). For the KA, the KA$_{\text{ox}}$ and the KA$_{\text{dith}}$ samples the broad signal was different with respect to its spectroscopic parameters, $g_{\text{eff}}$ and $\delta B$. The untreated sample revealed a $g_{\text{eff}}$ of 2.119. A shift to higher value, that is, to lower applied field, was observed in chemically treated samples. For KA$_{\text{ox}}, g_{\text{eff}} = 2.145$ and for KA$_{\text{dith}}, g_{\text{eff}} = 2.29$ were determined. For KA and KA$_{\text{ox}}, \delta B$ was similar with values of 122 and 119 mT, respectively. In contrast, a higher $\delta B$ of 161 mT was found for KA$_{\text{dith}}$. Replicate measurements with removal of the samples from the spectrometer cavity between the runs led to slight changes in the FMR parameters. The variation was most pronounced in KA$_{\text{dith}}$.

A similar effect was found when rotating the sample in a cavity. Fig. 6 shows the measured $g_{\text{eff}}$ versus rotation angle for one replicate run of each sample. The replicates for each sample differed slightly, but the relative differences between the samples were the same in all cases. The $g_{\text{eff}}$-values revealed for all the three samples an angular dependence with no overlapping values between the samples. Sample KA$_{\text{dith}}$ exhibited the most pronounced angular dependence. In this sample, $g_{\text{eff}}$ was characterized by a mean value of 2.27 and $\sigma = 0.05$. A periodicity of 180° was found (Fig. 6). In the two other samples, the angular dependence was weak. For KA$_{\text{ox}}$ the mean value for $g_{\text{eff}}$ was 2.142 with $\sigma = 0.004$, whereas in KA, the mean value for $g_{\text{eff}}$ was 2.119 with $\sigma = 0.008$. Furthermore, an angular dependence was also found for $\delta B$ (Fig. 7). In KA the mean $\delta B$ was 122 mT with $\sigma$ of 4.6 mT, while in KA$_{\text{ox}}, \delta B$ was 121 mT with $\sigma$ of 2.2 mT. For KA$_{\text{dith}}$ $\delta B$ had a mean value of 164 mT with $\sigma$ of 20 mT. Upon rotation the $\delta B$ was characterized by broad maxima and minima and the occurrence of a shoulder as additional feature in the EPR spectra, indicating a superimposed signal (Fig. 7b). This feature appeared simultaneously with the increase of the linewidth and was most pronounced where the $\delta B$ reached its maximum (Fig. 7).

The KA spectra could be reasonably fitted with a single Lorentzian line, whereas the best fitting for chemically treated samples was obtained by the linear combination of Lorentzian and Gaussian lines. Sample KA$_{\text{ox}}$ could be fitted well by using two

Figure 1. Changes in $\chi$ upon heating and subsequent cooling of sample KA.

Figure 2. Magnetization versus applied field plot of KA and KA$_{\text{acid}}$.

Semi-quantitative EDX analysis revealed varying substitution by minor amounts of cations, mainly Ti. A few magnetite grains contained an amount of substitution which was below the detection limit of the EDX analyses.

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components with the same $g_{\text{eff}}$ and similar $\delta B$. For the spectra of KA_dith revealing the additional shoulder, the best fit was achieved by the combination of the components with different $\delta B$ as well as $g_{\text{eff}}$ (Fig. 8).

The EPR spectra at low temperature exhibited the expected increase in intensity for the paramagnetic signal and a slight decrease for the broad FMR (Fig. 9). The lineshape of KA and KA_ox showed no significant change whereas $\delta B$ increased at $T < 100$ K (Fig. 9). Between 50 and 10 K the $\delta B$ of KA and KA_ox revealed a pronounced increase of about 60 and 30 per cent, respectively (Fig. 10). The $\delta B$ of KA_dith exhibited a different behaviour. It increased slightly upon cooling to 120 K, then $\delta B$ almost doubled between 120 and 100 K, followed by a steady increase down to 10 K. Due to the superimposed paramagnetic signal, $g_{\text{eff}}$ for the FMR at low temperatures could not be determined exactly from the spectra.

4 DISCUSSION

The hysteresis loop of the untreated sample shows that the remanent magnetic properties are dominated by a low coercivity phase. The $\chi$ versus temperature measurement shows that this phase has a $T_c$ of about 850 K, which is indicative of magnetite. The semi-quantitative EDX analysis shows the presence of minor amounts of Ti in the magnetite grains. The Ti content, however, has to be relatively low, because it is not documented in a significant lowering of $T_c$ (e.g. Dunlop & Ozdemir 1997). The bump in $\chi$ at about 570 K is probably due to the conversion of ferric hydroxides into an oxide, most likely hematite. The lack of clear evidence for a Neel temperature of hematite in $\chi$ at $T > 380$ K points to a very low concentration of this ferric oxide in the heated sample. Upon cooling the increase in $\chi$ indicates the generation of additional magnetite as a conversion product most likely of clay minerals.

The magnetite particles in the soil samples resist chemical treatment with oxalate and CBD as indicated by the hysteretic properties. This is supported by the similar morphologies of the grains under the SEM (Fig. 4). Gehring et al. (1997) showed that oxalate extracts about 10 per cent of the total Fe in the sample. The oxalate extractable Fe can be attributed to poorly crystallized ferric oxides which do not significantly contribute to the magnetic remanence of the sample. The difference in $\chi$ between KA and KA_ox is smaller than the variability of $\chi$ in the original topsoil material ranging between 74 and $86 \times 10^{-6}$ m$^3$ kg$^{-1}$ (Gehring et al. 1997). With this in mind it is postulated that KA and KA_ox are similar with respect to their $\chi$. The CBD treatment is known to dissolve ferric oxides such as hematite and maghemite. Furthermore, CBD can also dissolve superparamagnetic (SP) and SD magnetite (Hunt et al. 1995; Van Oorschot & Dekkers 1999). In our soil sample, CBD extracts 25 per cent of the total Fe (Gehring et al. 1997). The fact, that CBD extracts more Fe than oxalate, indicates a significant content of crystalline ferric oxides that contribute to the remanence of the sample. This agrees well with the decrease in $\chi$ in the CBD treated sample, which is most likely due to the removal of maghemite, since no clear indication for hematite has been found in the soil samples. The $\chi$ values are low compared to the bulk susceptibility of magnetite and titanomagnetite found in the literature (e.g. Peters & Dekkers 2003). These authors reported average $\chi$ values of $67 \times 10^{-5}$ m$^3$ kg$^{-1}$ for magnetite, $42 \times 10^{-5}$ m$^3$ kg$^{-1}$ for titanomagnetite and $63 \times 10^{-5}$ m$^3$ kg$^{-1}$ for maghemite, which are about three orders of magnitude higher than the susceptibility of our soil sample. Therefore, assuming that susceptibility arises only from ferrimagnetic phases,
a concentration of magnetite/maghemite of approximately 0.1 wt per cent can be deduced for it.

The hysteretic properties (i.e. \( B_c \) and \( M_r/M_s \)) of KA, KA_ox, and KA_dith give clues to PSD magnetite, which is characterized by grain sizes in the range from 0.3 to approximately 20 \( \mu \)m (Dunlop & Özdemir 1997). Parameters indicative of PSD grains are often found in soils and sediments and can also be produced by a mixture of particles with a wide variation of grain sizes (Roberts et al. 2000).

Weakly diverging open contours in the FORC diagram as found for KA, KA_ox and KA_dith (Fig. 3) have been described for magnetite in PSD state (Carvallo et al. 2005). The spread along \( B_u \) in the FORC diagram has been denoted as reversal ridge and can be generated by SP phases (Roberts et al. 2000) or by MD particles (Pike et al. 2001). Since SP magnetite is ruled out (Gehring et al. 1997), the reversal ridge is most likely caused by MD particles. It follows that the magnetite grains in the soil can be considered as PSD to MD. This is in good agreement with the SEM observations which revealed magnetite grains of a size generally larger than 20 \( \mu \)m. The similarity of the FORC diagrams, however, provides clear evidence that the grain size distribution of magnetite has not changed during the chemical treatments.

The EPR spectra of all samples except KA-acid are dominated by a broad signal characteristic for FMR, but they differ in their EPR parameters \( g_{\text{eff}} \) and \( \delta B \). It is well known that magnetite and maghemite can produce strong FMR signals (e.g. Griscom 1984; Gehring & Karthein 1990). Other ferric oxides such as goethite and hematite can also cause a broad resonance (Angel & Vincent 1978; Gehring & Karthein 1989). Because the absorption intensity is proportional to the susceptibility, the EPR signal of these ferric oxides will be weak and hidden by the signal arising from magnetite/maghemite. Therefore, the removal of goethite and hematite cannot account for the observed spectral changes upon CBD
The observed changes could result from the dissolution of SD magnetite and maghemite or by partial chemical modification of MD or PSD magnetite. The similarity of the FORC diagrams before and after CBD treatment argues against significant changes of the magnetite properties, and thus in favour of the dissolution of the maghemite. The occurrence of a shoulder in the KA_dith FMR spectrum upon rotation suggests an additional directionally dependent absorption in the sample where maghemite has been removed. Schlömann & Zeender (1958) describe additional subsidiary peaks in polycrystalline ferrites occurring on the low-field side of the main resonance and explain them by strong magnetocrystalline anisotropy. Therefore, it is assumed that the shoulder found in the KA_dith spectra is caused by the magnetocrystalline anisotropy. With this in mind, a likely reason why the secondary absorption described above is clearly visible in the KA_dith only, is that maghemitization reduces or masks the magnetocrystalline anisotropy. With other words, there is a link between CBD treatment and the increase of the magnetocrystalline anisotropy. Considering the maghemitization as a topotactical transformation (Gallagher et al. 1968), the dissolution of maghemite increases the Fe^{II}/Fe^{III} ratio in the crystal structure. It is known that magnetocrystalline anisotropy energy in magnetite arises mainly from Fe^{II} (e.g. Dunlop & Özdemir 1997), and, therefore, the maghemitization, that is, oxidation of Fe^{II}, reduces it, suppressing the secondary EPR absorption in KA and KA_ox. The angular dependence of this absorption, which is clearly visible for KA_dith, was not considered by Schlömann & Zeender (1958). Despite this fact, the behaviour of the secondary peak can be explained based on their findings. These authors showed for homogeneous polycrystalline systems a dependence of the intensity of the subsidiary absorption from the magnitude of the anisotropy and its influence on the effective field. In such a system, one can assume a randomized distribution of the anisotropies of the individual grains, and thus, of the corresponding easy axes. In natural samples, however, the magnetite grains are highly diluted in the soil matrix. In our experimental design, these magnetite grains in untreated and chemically treated samples are fixed relative to each other. As a consequence, the distribution of the easy axes has a given preferential...
orientation which leads to the angular dependence of the secondary EPR absorption indicated by $g_{eff}$ (Fig. 6). The angles where $g_{eff}$ has its maxima can be assigned to the configuration where most of the easy axes have the lowest angular departure relative to the applied field (e.g. Bickford 1950).

The low temperature EPR data provide further evidence for the maghemitization of magnetite in our tropical soil. The jump in $\delta B$ for KA_dith between 100 and 120 K occurs in the same temperature range where the Verwey transition of magnetite has been observed. At this transition, the crystallographic structure changes from cubic to monoclinic (e.g. Muxworthy & McClelland 2000). Only a few low-temperature EPR data are available on natural magnetite (Gehring & Karthein 1989; Weiss et al. 2004; Ananou et al. 2005; Kopp et al. 2006). Weiss et al. (2004) used the increase in $\delta B$ of 30 per cent between RT and 77 K as an indication for a Verwey transition. The samples KA and KA_ox, which show no jump between 120 and 100 K exhibit an increase between RT and 80 K by about 50 per cent. Hence, the broadening in such a temperature range cannot be taken as an unambiguous evidence for a Verwey transition. Ananou et al. (2005) published a sequence of low temperature EPR data of a tephra containing magnetite/maghemite and another ferric oxide, in which no spectral changes between 120 and 100 K were found. The EPR data on synthetic magnetite reported in the literature show also no uniform picture of the Verwey transition. Different phenomena such as line broadening, loss of absorption or the appearance of new signals have been described associated with this transition (e.g. Bickford 1950; Kale et al. 2001; Stankowski et al. 2006). A jump in linewidth as found for KA_dith was observed in a zero field cooled Fe$_3$O$_4$ crystalline film (Kale et al. 2001). These authors explained this phenomenon by an increase in possible easy axes caused by the crystallographic changes from cubic to monoclinic. With this in mind, we assume that the observed jump in $\delta B$ indicates a Verwey transition in KA_dith. The absence of this jump in the spectra of the KA and KA_ox samples indicates a suppression of the Verwey transition. Such a suppression can be caused by either oxidation, that is, maghemitization, defect structures (Ozdemir et al. 1993), or by an increased Ti content (Kakol et al. 1992). The CBD treatment is known to dissolve maghemite (Van Oorschot & Dekkers 1999), but leaves larger magnetite grains ($>20 \mu m$) mainly unaffected (Hunt et al. 1995). Therefore, the visibility of the Verwey transition in KA_dith is most likely due to the removal of maghemite associated with the magnetite surface. Finally, such oxidation of the magnetite surfaces fits well in the depositional environment of a subtropical alluvial Vertisol which is characterized by relative fast weathering dynamics.

5 CONCLUSION

The combination of EPR spectroscopy and rock magnetic methods applied to untreated and chemically treated soil samples allowed to detect the maghemitization of magnetite. The rock magnetic methods identified maghemite as the only carrier of the remanent magnetization at RT. The FORC diagrams showed that the different chemical treatments did not change significantly the magnetic properties of the magnetite grains. The EPR spectral properties of the CBD treated sample, from which maghemite was extracted, provided an indirect proof for the maghemitization of the magnetite surface. The elimination of the surface oxidation by removing the oxidation product increased the magnetocrystalline anisotropy of the magnetite at RT as indicated by the angular dependence of $g_{eff}$ and $\delta B$. At low temperature, the jump in $\delta B$ between 100 and 120 K indicating a Verwey transition in the magnetite is also visible after the removal of maghemite. The presented experimental approach clearly demonstrates that EPR spectroscopy can be an additional tool to detect maghemitization in natural samples.

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REFERENCES


Gehring, A.U., & Karthein, R., 1989. ESR Study of Fe(III) and Cr(III) Hydroxides, Naturwissenschaften, 76, 172–173.


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