Single-domain chemical, thermochemical and thermal remanences in a basaltic rock
Ulrike Draeger, Michel Prévot, Thierry Poidras, Janna Riisager

To cite this version:

HAL Id: hal-00012641
https://hal.archives-ouvertes.fr/hal-00012641v4
Submitted on 11 Sep 2006

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Single-domain chemical, thermochemical and thermal remanences in a basaltic rock

Ulrike Draeger,* Michel Prévot, Thierry Poidras and Janna Riisager†
Laboratoire de Tectonophysique, CNRS-UM2, Université de Montpellier 2, 34095 Montpellier Cedex 5, France. E-mail: prevot@dstu.univ-montp2.fr

Accepted 2005 October 26. Received 2005 October 15; in original form 2005 May 4

SUMMARY
Tiny basaltic samples containing finely grained titanomagnetite with Curie temperature less than 100°C were heated in air in weak field (25–100 μT) at temperatures between 400°C and 560°C for times as long as 32 hr. Oxyexsolution of titanomagnetite resulted in the crystallization of interacting single domain particles with Curie point close to 540°C and the concomitant development of one of two types chemical remanence, depending upon thermal treatment: isothermal chemical remanence (CRM) or thermochemical remanence (TCRM), the latter acquired under the combined effects of chemical change and temperature decrease. CRM and TCRM acquired under various conditions were subjected to Thellier-type experiments. All these treatments were carried out using a vibrating sample thermomagnetometer allowing the continuous recording of magnetization and a very precise temperature control. All CRM–TRM and TCRM–TRM plots were found to be linear over almost the entire TRM blocking temperature range, whether pTRM checks are positive or not. An apparent strength of the acquisition field of CRM or TCRM could thus be obtained and divided by the actually applied field to obtain a ratio R, which is representative of the CRM/TRM or TCRM/TRM ratios over most of the unblocking/blocking temperature spectrum. For CRM, it is found that R is less than 1 and increases rapidly with acquisition temperature (0.36 ± 0.07 at 400°C, 0.66 ± 0.02 at 450°C, and 0.90 ± 0.02 at 500°C), in qualitative agreement with expressions derived from the theory of non-interacting single domain grains. Thus, very large underestimate of geomagnetic field palaeostrength can occur when a natural CRM is not recognized as such and is believed to be a natural TRM. Palaeointensity data obtained from geological material prone to the development of secondary minerals, as for example baked contacts and volcanic glasses have, therefore, to be considered with caution. In an attempt to mimic deuteric oxyexsolution in cooling magma, TCRM was imparted from 560°C to 400°C at a cooling rate of 0.1°C/min. In contrast to the CRM case, R is found to slightly exceed 1 (1.11 ± 0.05). According to theoretical considerations, this finding suggests that in practice TCRM approximates to TRM.

Key words: basalt, geomagnetism, palaeointensity, palaeomagnetism, rock magnetism, titanomagnetite.

1 INTRODUCTION
It is a common feature in basaltic rocks that primary titanium-rich titanomagnetite, after crystallizing at high temperature subsequently undergoes some oxyexsolution during magma cooling. The highly magnetic mineral formed during this subsolidus chemical reaction is a very finely grained titanium-poor titanomagnetite. This is a magnetic carrier particularly appreciated in palaeomagnetism because of the high stability of its remanences versus field, temperature and time. Moreover, this mineral is generally magnetically and chemically stable when heated at moderate temperatures. For all these reasons, volcanic rocks carrying oxyxolved titanomagnetite are commonly preferred for studying the intensity of the Earth’s magnetic field over geological time.

Palaeointensity methods using volcanic rocks assume that NRM is a TRM. However, this fundamental assumption cannot be correct if oxyexsolution continues down to temperatures less than the Curie temperature of titanium poor titanomagnetite, that is commonly found to be around 550°C. An extensive magnetic study of magnetic oxides in two slowly cooling Hawaiian lava lakes led Grommé
et al. (1969) to the conclusion that titanomagnetite oxyexsolution in nature occurs between 850 °C and 500 °C, and even possibly down to 300 °C. Typically, most of the blocking temperatures of oxyexsolved titanomagnetite are above 500 °C. Therefore, there is a distinct possibility that the primary remanence acquired by some volcanic rocks is a thermochemical remanent magnetization (TCRM) rather than a TRM (Geissman & van der Voo 1980). In the present paper, we define a TCRM as a magnetization acquired under the combined influence of chemical change and temperature decrease. In contrast, we will use the expression ‘chemical remanent magnetization’ (CRM) for referring to a magnetization acquired as a result of chemical changes occurring at any constant temperature.

The objective of the work reported in this paper is to compare some characteristics of TCRM, CRM and TRM imparted to basaltic rock samples in the laboratory. Previous works in that field are restricted to the CRM case. The pioneer experimental studies of Haigh (1958) and Kobayashi (1959) showed that the intensity of CRM acquired by magnetite while it forms at 300 °C is much weaker than that of TRM. Later on, Stacey & Banerjee (1974) demonstrated theoretically the general validity of that conclusion for non-interacting single domain particles. They showed that the CRM/TRM ratio depends on the ratio of the blocking temperatures of CRM and TRM and reaches the maximum value of 1 when these temperatures are equal. Most of subsequent experimental data obtained either with synthetic minerals (Pucher 1969; Ik Gie & Biquand 1988; Stokking & Tauxe 1990) or rocks (Hoye & Evans 1975; Nguyen & Pechersky 1985) confirmed that low-field CRM magnetite acquired in the 300–500 °C range is generally largely inferior to TRM. However, Pucher (1969) and Kellogg et al. (1970) reported some cases in which CRM is equal to TRM. Obviously, the fact that CRM and TRM acquired in the same field can have largely different intensities may be the source of large error in paleointensity determination when a natural CRM is not recognized as such. Unfortunately, CRM and TRM cannot be easily distinguished from each other using usual palaeomagnetic techniques. According to Kobayashi (1959) and Pucher (1969) CRM and TRM generally have similar stability with respect to both AF and thermal demagnetization. Nevertheless, McClelland (1996) suggested from theoretical considerations that CRM and TRM carried by non-interacting SD grains can be distinguished from each other on the basis of Thellier paleointensity experiments.

Our samples were selected from a rock containing very fine titanomagnetite particles, with the hope that the magnetic carriers exsolved during CRM or TCRM acquisition will be within the single domain range. This turned out to be the case, which allows an evaluation of our experimental data in light of SD theories. In this paper we will use the Néel theory (1949) of single domain grains. Although some aspects of this theory can be challenged, as we will discuss in the present paper, this is to date the only theoretical approach allowing to deal in a quantitative manner with most aspects of remanent magnetism in rocks, in particular CRM and TCRM intensities. In order to evaluate the reliability of paleointensity determinations, our study was focussed on the comparison of intensity and thermal-unblocking/blocking of thermal (TRM) chemical (CRM) and thermochemical (TCRM) remanences. The main difficulty encountered in such studies is that, in contrast to deuterium oxyexsolution, oxyexsolution under laboratory conditions does not allow reaching a complete equilibrium of the newly formed phases with their oxidizing atmosphere. These exsolved phases are unstable during the subsequent heating needed for imparting a TRM. To try to evaluate the magnetic effects of the chemical changes occurring after CRM or TCRM acquisition, the TRM acquisition in the present work was carried out using the Thellier palaeointensity method.

2 EXPERIMENTAL PROCEDURE AND EQUIPMENT

2.1 Sample selection: palaeomagnetic and initial rock magnetic experiments

Several hand samples were taken from a very thin (about 15–20 cm thick) Quaternary basaltic dyke cutting a Jurassic limestone cliff close to the town of Carlenças (Southern France) along D908 (43.683° N, 3.228° E). Two hand samples (CAR4 and CAR5) were used for the present study. Seven small cores (12 mm in diameter) were drilled out of hand sample CAR4, and 22 out of hand sample CAR5. Each drill core was then cut into up to seven 7-mm-long specimens, labelled ‘a’ to ‘g’, where ‘a’ is the specimen close to the surface. Altogether about 120 specimens of about 2 g in weight, 12 mm in diameter and 7 mm in length were labelled and used for our experiments. For example, specimen CAR51d is from hand sample CAR5, core number 1, and is the fourth specimen from the surface. In addition two 1-inch diameter cores were drilled out of CAR5 in order to perform Thellier palaeointensity experiments.

The NRM of the samples, measured using a JR5 spinner magnetometer, was progressively demagnetized up to 200 mT using a laboratory made alternating field demagnetizer, and the median alternating destructive field (MDF) was determined. Low field (0.1 mT) thermomagnetic curves were obtained from some 40 specimens using either a Bartington susceptibility meter equipped with a furnace in which samples could be heated either in vacuum or in air or a Kappabridge in which specimens were heated in helium. The average Curie temperature of each magnetic phase was determined from the inflection point on the k-T curve (Prévôt et al. 1983). In complement, several samples were studied at the palaeomagnetic laboratory in Munich to measure hysteresis cycle at room temperature and thermal variation of saturation magnetization. Jurassic.

40 fresh looking specimens were finally selected for CRM and TCRM acquisition experiments on the basis of the following criteria: low Curie temperatures (between 40 °C and 80 °C), MDF between 30 and 50 mT, and position in the hand sample as remote as possible from the surface (no ‘a’ specimens were used).

2.2 Acquisition of chemical remanences

CRM and TCRM acquisition and measurement were carried out in air using a vibrating sample magnetometer (for short, VTM in the following) manufactured by Orion (Yaroslavia, Russia) and modified by one of us (TP) in respect to temperature control and magnetic signal sampling. The magnetic sensor is composed of two identical detection coils connected in opposition in order to cancel ambient field fluctuations with zero- or first-order gradients. This magnetometer is surrounded by two concentric shield screens made of Mumetal in order to try to cancel the ambient field at the centre of the magnetometer. Before the experiments reported here, the static residual field was checked and found to be less than 50 nT in the entire sample displacement space. The sample is alternatively moved from one detection coil to the other with a frequency of 13.7 Hz and an amplitude of 25.4 mm generating an electromotive force, which is filtered and pre-amplified before data acquisition by the computer. The magnetization is measured continuously, each registered data being sampled over 14 s. The non-inductive heater is
made of two oppositely wound resistor wires driven by a pulse width modulated (PWM) power supply. The base frequency of this PWM power supply is 3740 Hz. The control of the temperature is done by an Eurotherm 808 controller and the temperature is measured with a thermocouple S placed in a little hole in the sample. Our software makes it possible to define three successive thermal stages, each with their proper heating or cooling rate and applied dc field, through the serial port of the computer connected to an Eurotherm controller. A solenoid located between the detection coils and the furnace enables us to apply a continuous inducing field up to 1.2 mT upon the sample during the experiment.

In our experiments, the thermal cycle for CRM acquisition was carried out in air and consisted of three stages:

1. stage 1: rapid heating (10°C/min) from \( T_0 \) (room temperature) to \( T_r \) (temperature of CRM acquisition); the same field was applied during stages 1 and 2 in order not to impart a high temperature IRM to the sample at the beginning of stage 2;
2. stage 2: constant temperature, typically maintained (within 1°C) for 32 hr in a 100 \( \mu \)T dc field;
3. stage 3: rapid cooling (10°C/min) in zero field from \( T_r \) to \( T_0 \).

Approximately 30 CRM acquisition experiments were carried out, most of them at \( T_r = 400 \)°C, and a few at 450°C or 500°C. A few experiments were carried out in steady fields of 25 or 50 \( \mu \)T. Two experiments were made under a field of 100 \( \mu \)T, which was regularly turned off and on for a few minutes. These latter experiments were used to determine the ratio of remanent to induced magnetization, and for estimating short-term magnetic viscosity. These samples were not used to estimate the CRM/TRM ratio.

In order to monitor changes in magnetic mineralogy during CRM experiments, a distinct set of 13 fresh tiny specimens was heated to 400°C, each for a different duration distributed between 1 and 32 hr, using the same VTM equipment and procedure at Montpellier laboratory. Subsequently, hysteresis curves at 400°C were measured on these specimens at the Saint Maur geomagnetic laboratory using a laboratory-made translation inductometer (Bina & Daly 1994) including a furnace and a field coil providing fields up to 0.2 T. The maximum field we used for these hysteresis curves was up to 190 mT.

The thermal cycle for TCRM acquisition, also carried out in air, consisted of the three following stages:

1. stage 1: rapid heating (10°C/min) in zero field from \( T_0 \) (room temperature) to 560°C, a temperature which exceeds the Curie point of oxidized samples by some 20°C (Section 4.1);
2. stage 2: slow cooling (0.1°C/min) in a 100 \( \mu \)T constant field down to 400°C;
3. stage 3: rapid cooling (10°C/min) in the same field from \( T_r \) to \( T_0 \).

Three tiny specimens were subjected to this treatment. For four others the field was alternatively turned on and off every 47 or 68 s.

2.3 Thellier experiments on chemical remanences

The Thellier experiments (Thellier & Thellier 1959) were carried out in air using our modified VTM. TRM and CRM (or TCRM) were directed along the length of the tiny specimen, all with the same sense. We used the Coe variant (Coe 1967) of the Thellier method, and we carried out pTRM checks (Thellier & Thellier 1959) at every temperature steps. For each sample, the applied field was the same as for CRM or TCRM acquisition. For each Thellier run, all remanence measurements were carried out at the same temperature. For CRM, this measurement temperature was the same as the temperature of CRM acquisition (400°C, 450°C or 500°C). For TCRM, the measurement temperature was the lowest temperature of the interval of TCRM acquisition (400°C). Thus, no magnetic phase with Curie temperature less than 400°–500°C can possibly be involved in Thellier experiments.

Due to software constraints, temperature steps had to be constant for each Thellier run. The first experiments carried out with CRM involved temperature steps of 20°C. For better resolution of the pseudo-palaeointensity data, these steps were subsequently reduced to 10°C, and finally to 4°C for the latest experiments. In parallel, the first temperature step was increased from 410°C to 490°C, since no significant CRM decrease or pTRM acquisition had been observed within the 400°–490°C interval. For the Thellier experiments on TCRM, we used temperature steps of 10°C from 410°C to 560°C (or 580°C). The rate of increase or decrease of temperature employed was 4°C/min. When the desired temperature was reached it was held for 5 min. The measured temperature drift never exceeded 1°C.

2.4 Final rock magnetic experiments

The objectives of these experiments were to characterize the initial magnetic phase, the phases formed during CRM acquisition, and subsequent changes possibly induced by repetitive heatings at higher temperatures. The following experiments were made either at the Montpellier laboratory, at the Saint Maur laboratory, at CEREGE (CNRS and Aix-Marseille University) or at the Institute for Rock Magnetism (IRM) at Minneapolis University, MN:

1. low and high field thermomagnetic curves (Montpellier and IRM). Low field experiments carried out at both places used a Kappabridge (field intensity 0.4 mT). High field experiments used the MicroVSM (maximum field intensity 0.5 T) at the IRM.
2. room temperature hysteresis measurements (Saint Maur, IRM and CEREGE). Bulk hysteresis data were obtained either at Saint Maur using the laboratory-made translation inductometer described above or at the IRM using a MicroVSM (maximum field intensity 1 T). Differential hysteresis measurements such as first-order reversal curves (FORC) were obtained at CEREGE from measurements by a Micromag 3900 VSM.
3. thermal variation of hysteresis characteristics (Saint Maur and IRM). Same equipment as described in item 2.
4. AF demagnetization of pTRM (520°C, 515°C) (Montpellier and Saint Maur). This pTRM was imparted using the Orion VTM, and then demagnetized at room temperature using similar laboratory-made alternating field demagnetizers.

3 EXPERIMENTAL RESULTS

3.1 Starting material

The low-field thermomagnetic curves of unheated material (e.g. CAR517g in Fig. 1) exhibit Curie temperatures varying from 35°C to 80°C. This indicates titanomagnetite with high titanium content (x between 0.7 and 0.8). Hysteresis parameters plot in the central area of the PSD domain for Ti-rich titanomagnetite (Fig. 2, samples CAR517f, CAR517g). Microscopy study of polished sections revealed a bimodal grain-size distribution of titanomagnetite with modes around 2 \( \mu \)m and 20 \( \mu \)m. No ilmenite was observed,
neither as discrete crystals nor as exsolution within titanomagnetite crystals.

The two 1-inch cores (CAR51B and CAR52C) were studied by the Thellier stepwise heating method. They do not exhibit any change in NRM direction during heating throughout the investigated temperature range. PTRM checks are positive up to 175 °C and the NRM–TRM plot is linear up to 190 °C (Fig. 3). The NRM fraction is 0.7 and the quality factor \( q \) (Coe et al. 1978) is close to 25. Such long linear NRM–TRM plots obtained from high titanium titanomagnetite are usually typical of very fine grains (Prévôt et al. 1983).

To better characterize the micromagnetic structure of magnetic carriers, both in fresh and in heated materials, two kinds of differential hysteresis analyses were carried out: FORC and differential IRM measurements. FORC diagrams (Pike et al. 1999; Roberts et al. 2000) represent the density distribution of irreversible differential magnetic moments in function of two fields mathematically defined (Roberts et al. 2000), and usually labelled \( H_c \) and \( H_u \). For a set of randomly oriented identical uniaxial single-domain particles with axial coercive force \( H_c \), the average \( H_c \) value obtained from FORC should approach 0.48 \( H_c \), the theoretical value of bulk coercive force, and \( H_u \) should be a local interaction field (Néel 1954). This interpretation is reasonable agreement with experimental FORC diagrams measured from well-characterized SD materials (Pike et al. 1999; Carvallo et al. 2004) or computed from various SD models (Pike et al. 1999; Carvallo et al. 2003; Muxworthy et al. 2004).

The interpretation of FORC diagrams provided by MD and PSD grains is more uncertain. As first proposed by Néel (1942), hysteresis due to wall movement can be formally treated as resulting from the combination of various types of elementary hysteresis cycles, as those represented in a Preisach-Néel diagram. However, this is a
normal magnetization curves to consider with caution. In particular, FORC diagrams obtained for MD grains reveal a more complex pattern than those which can be computed from simple MD models (Pike et al. 2001b). Therefore, the physical interpretation of the FORC coordinates remains unclear. Empirically, judging from the diagrams obtained by Roberts et al. (2000), rocks containing PSD and MD particles provide somewhat similar FORC diagrams characterized by a density of differential moments which is maximum at the origin, and a similar distribution of both Hu’s and Hc’s.

Magnetic after-effect (magnetic viscosity), which can affect all grain sizes from MD to small SD, is the source of some artefacts in FORC diagrams, as a secondary peak in moment distribution near the origin (Pike et al. 2001a) or, in our opinion, anomalies in moment distribution in the neighbouring of the diagonal of the inferior quadrant of FORC diagrams (as in the middle diagram of Fig. 4). We think that the latter anomalies are probably due to the fact that this diagonal is the locus where the remagnetizing field Hb changes sign during the acquisition of FORC data.

The FORC diagrams of fresh specimen CAR521d (upper two diagrams on Fig. 4) look intermediate between those of SD and PSD/MD grains. On one hand, the peak of moment distribution is located at the origin, on the other hand Hc distribution reaches values exceeding 150 mT while interaction fields are smaller by more than 1 order of magnitude. The weakness of interaction fields is independently corroborated by the small values of differential IRM (Petrovsky et al. 1993; Wehland et al. 2006). Differential IRM is the difference between the IRM measured in back field H after saturation in the opposite sense and the IRM value expected from Wolfarth’s (1958) relations between IRM acquisition and SIRM d.c. demagnetization for non-interacting SD grains. Non-zero differential remanence is attributed to interaction fields. Positive (negative) values indicate a net-magnetizing (net-demagnetizing) effect of magnetic interactions. For specimen CAR521d, differential IRM, particularly scattered in low back fields, remains close to zero for any applied field H (Fig. 5). Altogether, bulk hysteresis and differential hysteresis data suggest that the magnetic configuration of the fresh material is mostly PSD.

3.2 Chemical remanence acquisition

The VTM provides a continuously record of total magnetization J changes at temperature as experiment proceeds. When exposed to a non-zero field, the measured magnetization is the sum of a remanent and an induced component. The latter, calculated from field on/field off experiments, was found to represent only 2–3 per cent of the total magnetization for acquisition temperatures up to 450 °C, and 5–9 per cent for higher temperatures (acquisition of CRM(500 °C) and TCRM). Let us note that the percentages given above are approximately overestimated by a factor of 2 since the applied field is off during half of the duration of the CRM or TCRM acquisition process. The viscosity of remanence (viscosity index; Prévot 1981) measured over a few minutes, was found to be lower than 1 per cent. Thus J measurements provide a very good approximation of Jcrm or Jocrm, even though a small high-temperature VRM component is inevitably included in these measurements.

During the first stage of the thermal cycle of CRM acquisition (rapid heating to Tr under a constant field), J is found to decrease rapidly to be reduced to a few per cent of the room temperature magnetization at temperatures around 150–200 °C (Fig. 6a). Beyond 200 °C a small progressive increase is observed. This increase in J reflects essentially an increase in susceptibility related to the onset of chemical changes around 200 °C: field on/field off experiments indicate that at any temperature between 200 and 400 °C, J is essentially an induced magnetization. At the onset of the temperature dwell at 400 °C, only 25 per cent of J is a remanent magnetization. Thus the CRM acquired during heating from 200 °C to 400 °C is

Figure 2. Day diagram (Jrs/Js versus Hcr/Hc) at room temperature for two fresh samples (CAR517f and g, full circles) and sample CAR518e (full square), the latter previously heated at 400 °C for 32 hr in order to acquire a CRM. Dashed lines indicates the approximate location of the boundaries of SD, PSD and MD domains for magnetite (Day 1977; Dunlop 2002). As the upper limit of the Hcr/Hc ratio for PSD grains varies with titanium content, this limit is indicated by a dash-and-dot line for composition TM60.

Figure 3. Example of NRM–TRM diagram for fresh material (sample CAR51B). Linearly extrapolated initial NRM is 5.40 × 10⁻³ Am² kg⁻¹. Linearly extrapolated final TRM is 5.73 × 10⁻³ Am² kg⁻¹. Laboratory field was 40 μT. Fa is the estimated palaeofield magnitude and q the quality factor (Coe et al. 1978).
Chemical and thermal remanences

Figure 4. FORC diagrams for fresh sample CAR521d (upper two diagrams, the second corresponding to more detailed measurements) and sample CAR57b, the latter previously heated (first for 32 hr at 400 °C for CRM acquisition, then for Thellier and hysteresis measurements at temperatures between 400 °C and 580 °C). The physical meaning of fields $H_c$ and $H_u$ is discussed in Section 3.1.

Figure 5. Differential IRM versus back dc field for chips of fresh sample CAR521d and previously heated sample CAR57b (see caption of Fig. 4 for description of previous thermal treatment).

extremely small compared to the final CRM (Figs 6a and b; note that the $J$ scale differs by one order of magnitude between these two diagrams). CRM develops rapidly as soon as temperature is stabilized at 400 °C. Field on/field off experiments indicate that only a few minutes after the onset of this temperature dwell, CRM reaches already 75 per cent of $J$.

Typically, CRM growth follows approximately a log $t$ law up to the end of the constant temperature stage (32 hr) (Figs 6b and 7a). This simple behaviour was also reported by Gapeev et al. (1991) for CRM resulting from low-temperature oxidation of nearly single-domain magnetite. In fact, more detailed calculations show that in our case CRM acquisition passes through a maximum 1 hr after the beginning of the constant temperature stage (Fig. 7b). An approximately log $t$ acquisition law corresponds the simplest behaviour, which can be derived from the Néel (1949) relaxation theory of single-domain particles. However, for about one-third of our CRM acquisition experiments, a marked $S$-shape was observed in $J$ as a function of log $t$. These samples were not studied any longer.

Additional experiments were carried out on a separate set of 8 specimens that were heated at 400 °C, each of them for a distinct duration between 2 and 32 hr. When normalized to the same inducing field, CRM intensity is found to reach that of NRM in less than 2 hr.

The remanence of samples meant for TCRM experiments decays rapidly during heating to 560 °C in zero field (Fig. 8a). In this diagram, the change in sign around 250 °C is an instrumental artefact due to a progressive drifting of the baseline at the high sensitivity required for these measurements. During the subsequent very slow cooling in a 100 μT field, TCRM acquisition starts only around 540 °C (Fig. 8b). The interpretation of the shape of the TCRM acquisition curve is delicate because of concomitant changes in temperature and time. A tentative calculation of the TCRM blocking curve will be discussed in Section 4.3.
Figure 6. Thermal demagnetization curve of residual NRM (after AF demagnetization) (a), and curve of CRM acquisition versus time at 400 °C (b), both being measured in field $h = 100 \mu T$, for sample CAR56d. In (b) the origin of the time axis is the beginning of the constant temperature stage.

3.3 Thellier experiments on chemical remanences

Examples of CRM–TRM diagrams are given in Fig. 9 for CRMs acquired at 400 °C or 500 °C, and TCRM–TRM diagrams are shown in Fig. 10. The data for all samples are listed in Table 1 following the canon of presentation of Thellier palaeointensity data (Coe et al. 1978). An additional column gives $R$, the ratio of the apparent palaeointensity to the actual acquisition field. For this calculation, it was assumed that CRM is proportional to applied field in the investigated range (25–100 $\mu T$).

All plots are almost perfectly linear over large CRM fractions $f$ (from 0.61 to 0.95). $f$ is essentially dependent on the magnitude of the low-temperature CRM decrease (from 400 °C to, say, 510 °C) in which (CRM, TRM) couples do not fall in line with higher temperature data. As we will see below (Section 5), the unblocking/blocking processes are ineffectual in that low-temperature interval. The changes in CRM and TRM, which are depicted on the diagrams of Figs 9 and 10 in that interval result essentially from mineralogical transformations. Moreover, those changes are dependent on the mode of heating from 400 °C to 490 °C (Fig. 16 below).

The quality factor $q$ is largely controlled by the number of heating steps. Considering the data obtained using 4 °C temperature intervals, $q$ is about 30 for TCRM while it varies from 12 to 64 for CRM. Linearity of the diagrams derived from Thellier experiments is sometimes considered as an argument in favour of the absence of magnetic alteration during experiments (e.g. Stokking & Tauxe 1990). Obviously, this is not necessarily the case here. If we consider the four best-documented data set for CRM, obtained with 4 °C heating steps, only two samples yield positive $pTRM$ checks (like sample CAR56d in Fig. 9). Checks are fairly good for the second (sample CAR55e, Fig. 9) but they are clearly negative for the last one (sample CAR51d, Fig. 9): all checks carried out in the TRM unblocking interval indicate a significant increase in $pTRM$ at each temperature step. The three samples carrying a TCRM (Fig. 10) behave rather similarly to the last CRM-carrying sample. The origin and the possible effect on $R$ of the changes occurring during Thellier experiments, below and within the temperature interval in which plots are linear, will be examined in Section 5.

In practice, the $R$ ratio listed in Table 1 is a measure of the CRM/TRM (or TCRM/TRM) strength ratio over some 90 per cent of the TRM blocking range (Figs 9 and 10). For CRM’s, $R$ is approximately constant at a given temperature of acquisition $Tr$. For the best documented case ($Tr = 400 °C$) $R = 0.36 \pm 0.07$ for six determinations (in this paper, ± refers to standard deviation). Within the investigated range of $Tr$ (400°–500 °C) $R$ is always less than 1 and increases almost linearly with acquisition temperature (Fig. 11). In contrast, $R$ calculated from TCRM’s is slightly but systematically larger than 1.

© 2006 The Authors, GJI, 166, 12–32
Journal compilation © 2006 RAS
showed that phases 2 and 3 are not always distinct from each other
high T phase 3. Other low field experiments on different samples
heated above 400

magnetite. When samples having acquired a CRM are repetitively
±
phase 1, 450
440
diminishes while phase 1 grows, it does not unmix at high temper-

In Fig. 1 we can compare the thermomagnetic curves, both in
low and high-field, for a fresh sample (CAR517g) and a sample
previously heated in air at 400°C, each for a different duration between 1 and 32 hr, using the VTM
equipment at Montpellier laboratory (Section 2.2). The data ob-
tained from the hysteresis measurements at 400°C, subsequently
carried out at Saint Maur laboratory, are listed in Table 2 and shown
in Fig. 13. Although each data point corresponds to a distinct sam-
ple, the magnetic homogeneity of our sample set (see Section 2.1 for
selection criteria) is such that coherent trends versus heating time
are documented for all hysteresis data (Jr/Js, Hcr and Hcs/Js) with
the only exception of Hcr/Hc, which remains stationary. In spite of
the small irregularities due to the combination of data from distinct samples, three main stages can be readily distinguished during the
CRM acquisition process:

1. stage 1 (from 1 to 2 hr): large increase in both induced and
remanent saturation magnetizations without noticeable changes in
coeercivity, Jr/Js or Hcs/Hc (see Fig. 2 for comparison with fresh
material);

2. stage 2 (from 2 to 4 hr): steep increase in Jr/Js ratio from
0.30 to 0.40–0.45 accompanied by an increase in saturation magnet-
izations and coercive forces;

3. stage 3 (beyond 4 hr): saturation magnetizations continue
to increase while both Jr/Js ratio and coercivity remain almost
constant.

In addition, other experiments carried out on another set of spec-
imens (Section 3.2) showed that the median destructive AF of CRM
acquired at 400°C increases with time. Comparable to that of NRM
for t = 2 hr (36 mT), the MDF of CRM reaches 50–60 mT for
heating times between 20 and 30 hr.

Stage 1 may correspond to a monophase oxidation producing metastable titanomaghemite (Goss 1988) with progressively in-
creasing oxidation degree and Curie point. The crucial step in the
mineralogical process described above is the decrease in magnetic
grain size, which occurs during stage 2. This reduction is expected in
cases of exsolution, whether it occurs by spinodal decomposition or
nucleation and grow (Fig. 14). The brevity of this stage attests to the
simultaneity of mineralogical reactions affecting magnetic grains
within the whole specimen. After this stage, no significant change in
magnetic grain size is documented although some slight increase
in volume might be suspected for the largest heating times. The
range of the Jr/Js ratio during stage 3 is 0.40–0.45. These values
are only slightly smaller than the theoretically value of 0.5 expected
for SD grains with shape (and therefore uniaxial) anisotropy. Let us
remember that, in order to avoid any contribution from the magnetic
phases not involved in the CRM acquisition process, this ratio was
measured at 400°C, not at Tg. For reasons which remain unclear,
Jr/Js generally tends to diminish with temperature. This decrease
is about 20 per cent for naturally oxyexsolved titanomaghemite heated
to 400°C (Dunlop 1987). The Jr/Js ratio found for those samples at
that temperature seems therefore indicative of a truly SD magnetic

Figure 8. Demagnetization curve (a) of residual NRM (after AF demag-
netization) and TCRM acquisition (b) for sample CAR56b. In diagram (b)
time zero is set at the beginning of the steadily decreasing temperature stage. The progressive shift of the baseline visible in diagram (a) is due to the high
sensitivity required for measuring the very small residual NRM.

4 PROCESS OF CHEMICAL
REMANENCE ACQUISITION

4.1 Mineral process

In Fig. 1 we can compare the thermomagnetic curves, both in
low and high-field, for a fresh sample (CAR517g) and a sample
previously heated in air at 400°C for 32 hr (CAR518e). As usual,
low field curves are much more discriminative for distinguishing
coexisting magnetic phases. After CRM acquisition, the magnetic
phase present in the fresh material has almost entirely disappeared.
Instead, three magnetic phases with high Curie temperature are ob-
erved. Their respective average Curie temperatures determined by
the method of Prévot et al. (1983) are 522 ± 11°C for high T
phase 1, 450 ± 10°C for high T phase 2, and 390 ± 30°C for
high T phase 3. Other low field experiments on different samples
showed that phases 2 and 3 are not always distinct from each other
and exhibit somewhat variable Tc, although they remain in the range
350°–500°C. Such phases may be either titanomaghemite or titano-
magnetite. When samples having acquired a CRM are repetitively
heated above 400°C, an intermediate phase with Curie point around
440°C is often observed (Fig. 12). Although this phase progressively
diminishes while phase 1 grows, it does not unmix at high temper-

ture as expected for titanomaghemite. Thus, such intermediate Tc
phases correspond probably to titanomagnetite with x between 0.2
and 0.4. Phase 1, that is favoured by high temperature treatments,
seems to correspond to the dominant near-stoichiometric magnetite
found in intergrowths resulting from inversion of titanomaghemite
(Tucker & O’Reilly 1980; Özdemir 1987). Combining the data ob-
tained from different samples subjected to various high temperature
 treatments, we estimate the final Curie temperature to be approxi-
mately 540°C. This is compatible with a stoichiometric titanomagne-
tite with x = 0.06 and JS0 = 421 kA m−1 (Kawai 1956).

Progressive magnetic mineral changes during CRM acquisition
were monitored using a set of 13 fresh specimens heated to 400°C,
each for a different duration between 1 and 32 hr, using the VTM
equipment at Montpellier laboratory (Section 2.2). The data ob-
tained from the hysteresis measurements at 400°C, subsequently
carried out at Saint Maur laboratory, are listed in Table 2 and shown
in Fig. 13. Although each data point corresponds to a distinct sam-
ple, the magnetic homogeneity of our sample set (see Section 2.1 for
selection criteria) is such that coherent trends versus heating time
are documented for all hysteresis data (Jr/Js, Hcr and Hcs/Js) with
the only exception of Hcr/Hc, which remains stationary. In spite of
the small irregularities due to the combination of data from distinct samples, three main stages can be readily distinguished during the
CRM acquisition process:

1. stage 1 (from 1 to 2 hr): large increase in both induced and
remanent saturation magnetizations without noticeable changes in
coeercivity, Jr/Js or Hcs/Hc (see Fig. 2 for comparison with fresh
material);

2. stage 2 (from 2 to 4 hr): steep increase in Jr/Js ratio from
0.30 to 0.40–0.45 accompanied by an increase in saturation magnet-
izations and coercive forces;

3. stage 3 (beyond 4 hr): saturation magnetizations continue
to increase while both Jr/Js ratio and coercivity remain almost
constant.

In addition, other experiments carried out on another set of spec-
imens (Section 3.2) showed that the median destructive AF of CRM
acquired at 400°C increases with time. Comparable to that of NRM
for t = 2 hr (36 mT), the MDF of CRM reaches 50–60 mT for
heating times between 20 and 30 hr.

Stage 1 may correspond to a monophase oxidation producing metastable titanomaghemite (Goss 1988) with progressively in-
creasing oxidation degree and Curie point. The crucial step in the
mineralogical process described above is the decrease in magnetic
grain size, which occurs during stage 2. This reduction is expected in
cases of exsolution, whether it occurs by spinodal decomposition or
nucleation and grow (Fig. 14). The brevity of this stage attests to the
simultaneity of mineralogical reactions affecting magnetic grains
within the whole specimen. After this stage, no significant change in
magnetic grain size is documented although some slight increase
in volume might be suspected for the largest heating times. The
range of the Jr/Js ratio during stage 3 is 0.40–0.45. These values
are only slightly smaller than the theoretically value of 0.5 expected
for SD grains with shape (and therefore uniaxial) anisotropy. Let us
remember that, in order to avoid any contribution from the magnetic
phases not involved in the CRM acquisition process, this ratio was
measured at 400°C, not at Tg. For reasons which remain unclear,
Jr/Js generally tends to diminish with temperature. This decrease
is about 20 per cent for naturally oxyexsolved titanomaghemite heated
to 400°C (Dunlop 1987). The Jr/Js ratio found for those samples at
that temperature seems therefore indicative of a truly SD magnetic

© 2006 The Authors, GJI, 166, 12–32
Journal compilation © 2006 RAS
configuration. Beyond 10 hr or so the coercivity remains constant while saturation magnetizations continue to grow. As we will see below, CRM seems essentially carried by phase 1. Considering the large saturation magnetization of this phase, the microcoercive force $H_c$ of these SD particles is due to shape anisotropy. Assuming that magnetization reversal is coherent, we have then:

$$H_c = N_d I_s,$$

where $I_s$ is the spontaneous magnetization and $N_d$ the demagnetizing factor of the particle.

Thus, if the $J_s$ increase was due to an increase in spontaneous magnetization, then $H_c$ should grow with $J_s$. The absence of correlation between $H_c$ and $J_s$ suggests that during this final stage of CRM acquisition the increase in $J_s$ is mainly due to volume growth of the magnetized material (within the space limited by the ilmenite frame) while the $I_s$ value, and therefore the chemical composition, of the final magnetic phase remains unchanged. This suggestion is confirmed by low-field thermomagnetic experiments such as those shown in Fig. 12 that show that repetitive heating to high temperatures leads to a further development of the phase 1 mineral without
any significant increase in Curie temperature. Thus, the oxyexsolution process seems to result from nucleation and growth rather than from spinodal decomposition (Fig. 14).

The mineral process at work during TCRM acquisition from 560°C to 400°C is certainly the same as during isothermal CRM acquisition since this temperature range is well within the domain of oxyexsolution of titanium-rich titanomagnetite (O’Reilly 1984). There is, however, a distinct characteristic of TCRM acquisition when compared to CRM acquisition: all things being equal, the reaction rate diminishes drastically with time in the first case, while it is constant in the second case. For an activation energy of 240 kJmol\(^{-1}\), which is a typical value for cation diffusion process in minerals (Putnis 1992), the reaction rate is divided by 4000 as \(T\) decreases from 560°C to 400°C. From 560°C to 500°C, the reaction rate is already reduced by an order of magnitude. This has an important consequence: during the process of TCRM acquisition, most of the chemical changes are expected to occur within the highest temperature range.

4.2 Final size and shape of CRM carriers

Fig. 2 demonstrates that the CRM carriers are much smaller in size than the original high-titanium titanomagnetite crystals. However, these bulk measurements carried out at room temperature cannot be used to estimate the size of CRM carriers because of the presence in the heated sample of a remnant fraction of the original low Curie temperature phase (Fig. 1a). As pointed out in the previous section, the hysteresis measurements carried out at 400°C during CRM acquisition suggest that the final magnetic product of
the chemical changes is of SD size. This inference is clearly confirmed by the FORC diagram (lower diagram of Fig. 4) obtained at room temperature from sample CAR57b (previously heated for 32 hr at 400 °C, then used for various measurements at temperatures between 400 °C and 580 °C). This diagram exhibits only a single phase of high coercivity displaying typical SD characteristics. The peak in coercivity distribution is about 53 mT. Following Néel’s interpretation of the hysteresis cycle of SD particles oriented at random (1954), the coercive force of individual particles measured along the magnetization axis is 110 mT. A further important information from this diagram is the presence of large interaction fields, as expected within magnetite-ilmenite intergrowths (e.g. Davis & Evans 1976). The magnitude of the interaction field defined by the FWHM on the FORC peak (Muxworthy et al. 2004) is 45 mT. Moreover, the plot of the differential IRM versus the back applied field obtained from sample CAR57b shows a large negative peak (Fig. 5), which confirms the presence of interactions and suggests that they are dominantly negative.

A quantitative estimate of the size and axial coercivity of typical CRM carriers can be obtained from the SD theory by combining low-field thermoviscous blocking and AF unblocking experiments. This method, based on Néel (1949) theory of single-domain non-interacting particles, was used for thermoremanence (Dunlop & West 1969) and viscous remanence (Biquand et al. 1971; Prévot 1981). Bina & Prévot (1977) showed that hematite grain sizes calculated with this method are in reasonable agreement with direct size measurement from TEM observations. According to Dunlop & West (1969), the presence of magnetic interactions does not invalidate such calculations since we are dealing with low-field remanences. They argue that only the grains subjected to very small interaction fields could have their magnetic moments aligned by such applied fields. In practice, the method consists of solving simultaneously the blocking and unblocking equations.

Table 1. Results of Thellier experiments on CRM and TCRM (experimental procedure described in Section 2.3). \( T_r \): temperature of CRM or TCRM acquisition; \( \Delta T \): temperature interval used for apparent palaeointensity determination; \( N \): number of temperature steps used for determination; \( f \), \( g \), \( q \): CRM (or TCRM) fraction used for apparent palaeointensity determination, gap factor, and quality factor, respectively (Coe et al. 1978); \( H_{app} \): apparent palaeointensity value with associated error; \( R \): ratio of the apparent palaeointensity to the actual acquisition field.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Type</th>
<th>( T_r ), °C</th>
<th>Field, ( \mu T )</th>
<th>( \Delta T ), °C</th>
<th>( N )</th>
<th>( f )</th>
<th>( g )</th>
<th>( q )</th>
<th>( H_{app} \pm \sigma (H_{app}) ), ( \mu T )</th>
<th>( R \pm \sigma (R) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAR55e</td>
<td>CRM</td>
<td>400</td>
<td>25</td>
<td>514–538</td>
<td>7</td>
<td>0.65</td>
<td>0.80</td>
<td>12</td>
<td>7.1 ± 0.3</td>
<td>0.28 ± 0.01</td>
</tr>
<tr>
<td>CAR53d</td>
<td>CRM</td>
<td>400</td>
<td>50</td>
<td>510–540</td>
<td>4</td>
<td>0.82</td>
<td>0.58</td>
<td>8</td>
<td>14.9 ± 0.9</td>
<td>0.30 ± 0.02</td>
</tr>
<tr>
<td>CAR44a</td>
<td>CRM</td>
<td>400</td>
<td>100</td>
<td>500–560</td>
<td>4</td>
<td>0.95</td>
<td>0.50</td>
<td>5</td>
<td>43.5 ± 3.9</td>
<td>0.44 ± 0.04</td>
</tr>
<tr>
<td>CAR53a</td>
<td>CRM</td>
<td>400</td>
<td>100</td>
<td>500–540</td>
<td>5</td>
<td>0.91</td>
<td>0.62</td>
<td>12</td>
<td>42.2 ± 2.0</td>
<td>0.42 ± 0.02</td>
</tr>
<tr>
<td>CAR56d</td>
<td>CRM</td>
<td>400</td>
<td>100</td>
<td>514–534</td>
<td>6</td>
<td>0.61</td>
<td>0.78</td>
<td>16</td>
<td>39.5 ± 1.2</td>
<td>0.40 ± 0.01</td>
</tr>
<tr>
<td>CAR513d</td>
<td>CRM</td>
<td>400</td>
<td>100</td>
<td>506–534</td>
<td>8</td>
<td>0.83</td>
<td>0.82</td>
<td>64</td>
<td>32.6 ± 0.3</td>
<td>0.33 ± 0.01</td>
</tr>
<tr>
<td>CAR44b</td>
<td>CRM</td>
<td>450</td>
<td>100</td>
<td>500–560</td>
<td>4</td>
<td>0.85</td>
<td>0.37</td>
<td>10</td>
<td>66.1 ± 2.0</td>
<td>0.66 ± 0.02</td>
</tr>
<tr>
<td>CAR51d</td>
<td>CRM</td>
<td>500</td>
<td>100</td>
<td>514–542</td>
<td>8</td>
<td>0.75</td>
<td>0.85</td>
<td>43</td>
<td>88.9 ± 1.3</td>
<td>0.89 ± 0.01</td>
</tr>
<tr>
<td>CAR45b</td>
<td>TCRM</td>
<td>560–400</td>
<td>100</td>
<td>470–540</td>
<td>8</td>
<td>0.86</td>
<td>0.81</td>
<td>27</td>
<td>107.5 ± 2.7</td>
<td>1.08 ± 0.03</td>
</tr>
<tr>
<td>CAR56b</td>
<td>TCRM</td>
<td>560–400</td>
<td>100</td>
<td>470–540</td>
<td>8</td>
<td>0.88</td>
<td>0.79</td>
<td>33</td>
<td>108.9 ± 2.3</td>
<td>1.09 ± 0.02</td>
</tr>
<tr>
<td>CAR56c</td>
<td>TCRM</td>
<td>560–400</td>
<td>100</td>
<td>490–540</td>
<td>6</td>
<td>0.77</td>
<td>0.73</td>
<td>31</td>
<td>116.5 ± 2.1</td>
<td>1.17 ± 0.05</td>
</tr>
</tbody>
</table>

Figure 11. Mean CRM/TRM ratio versus the temperature of CRM acquisition. The experimental data (full circles) are from Table 1. Labels by each circle indicate the number of experimental data. Error bars correspond either to the apparent ‘palaeointensity’ error (at 450 °C and 500 °C) or the standard deviation of apparent ‘palaeointensity’ determinations (at 400 °C). Theoretical ratio and associated error bar were calculated from eq. (11) assuming a TRM blocking temperature of 525 ± 5 °C.

Figure 12. Two successive low-field thermomagnetic cycles in air for sample CAR513d. Previously, this sample had been heated at 400 °C for 32 hr to acquire a CRM and then subjected to Thellier experiments.
In low field the blocking expression at some decreasing temperature \( T > T_b \) is:

\[
vH_c^0 = \frac{2kT}{I_{S0}} \left( Q + \ln \tau_0 \right) F(T) F'(T),
\]

with \( F(T) = Is(T)/I_{S0} \) and \( F'(T) = Hc(T)/H_c^0 \).

The unblocking conditions under a large field \( H \) applied at room temperature \( T_b \) are:

\[
\frac{v(H_c^0 - H)^2}{H_c^0} = \frac{2kT_0 \left( Q + \ln \tau_0 \right)}{I_{S0}}.
\]

In these expressions \( k \) is Boltzman's constant, and \( v \), \( Is \) and \( Hc \) are volume, spontaneous magnetization and microscopic coercive force of particle, respectively. Following Néel (1949) notations, \( Q \) is a constant estimated to 22 for magnetite, \( \tau_0 \) is the blocking time (depending upon cooling rate) and \( \tau_u \) the unblocking time (depending upon AF frequency).

The thermal unblocking of CRM(400°C) as measured at 400°C is generally maximum around 520°C. As we simply wish to get an approximate estimate of size and coercivity of CRM carriers rather than a complete description of the density distribution of grains in the whole (\( Hc, v \)) diagram, a single pTRM was imparted to sample CAR57b, between 515 and 520°C, before FORC experiments. This pTRM was then progressively AF demagnetized at room temperature. The blocking temperature to be introduced in expression (2) is the mean temperature of the pTRM acquisition interval. Two distinct AF experiments provided the same median alternating destructive field (140 mT). This value of the unblocking field was introduced into expression (3). From the experimental blocking and unblocking procedures, we estimated that \( \tau_0 \cong 15 \) s and \( \tau_u \cong 6 \times 10^{-6} \) s. \( F(T) \) and \( F'(T) \) between \( T_0 \) and \( T \) could not be precisely measured on our samples because of the presence of several magnetic phases (e.g. Figs 1 and 12). Thus, those ratios were taken from previous experiments on a thermally stable basalt sample containing deuteritic oxyexsolved (titano)magnetite with Curie temperature close to that of phase 1 (Dunlop & West 1969; Dunlop 1987). This is probably the main source of uncertainty in our calculation. Other experimental sources of imprecision are the thermal instability of our samples and the fact that the temperature interval chosen for pTRM acquisition may correspond to grains that are not perfectly representative of the CRM carriers as a whole.

Our calculations give \( Hc^0 = 144 \) mT and \( v = 2.3 \times 10^{-3} \) \( \mu m^3 \). This microcoercive force is of the same order of magnitude as the mean axial coercive force calculated from the FORC diagram. As mentioned above, phase 1 is strongly magnetic and the coercivity can be assumed to be due to shape anisotropy. If coherent reversal of magnetic moment is assumed, then eq. (1) is valid. Given the dependence of \( Nd \) on the particle dimension ratio (see for example Nagata 1961), the calculated microcoercive force requires a dimension ratio close to 2. Thus, considering the calculated volume mentioned above, the particle length would be about 0.2 \( \mu m \).

The dimension ratio obtained from eq. (1) is possibly a minimum estimate. According to theoretical considerations, coherent rotation in magnetite probably occurs only in a restricted size range below 50 nm (Dunlop & Özdemir 1997). The mode of magnetization reversal of larger elongated particles, like the present CRM carriers, is believed to be incoherent (Moon & Merrill 1988; Enkin & Williams 1994), which in turn should reduce particle coercivity. Thus a larger dimension ratio of the CRM-carrying particles would be needed to account for the calculated axial coercivity. The reduction in coercivity depends both on the grain size and the mode of incoherent reversal (buckling, curling or fanning; see for example Dunlop & Özdemir 1997). Theoretical considerations suggest that prolate spheroid may present only two modes of incoherent reversal, curling and buckling, the latter one being restricted to extremely large aspect ratios (Aharony 2000), which are not realistic for natural crystals. Thus, the most probable non-coherent reversal mode of elongated magnetite particles is curling (Newell & Merrill 1999). Taking into account the volume of the CRM-carrying particles, curling would reduce their coercivity by a factor of 3 as compared to that given by eq. (1) (Luborsky 1961). To account for the observed CRM coercivity, the dimension ratio should be then increased to 10–20.

It must be noted, however, that we have no experimental evidence in favour of the occurrence of incoherent reversal of magnetization in the final, largest-size CRM carriers. Whichever is the mineralogical process giving birth to the magnetite-ilmenite intergrowth (Fig. 14), a growth in the size of the magnetic phase has to occur during CRM acquisition experiments. According to the theoretical inferences presented above, the reversal mode of the CRM-carrying particles should evolve with the heating time. Coherent rotation should predominate at the onset of our CRM acquisition experiments, the particle size being then just above the SP/SD threshold. Then curling should progressively becomes predominant since the final grain size is clearly above the SD/PSD threshold for equidimensional particles. Thus, the coercivity of the CRM carriers is expected to markedly

<table>
<thead>
<tr>
<th>Sample</th>
<th>Heating time at 400°C</th>
<th>( J_{is} ), ( 10^{-3} ) Am²/kg</th>
<th>( J_s ), ( 10^{-3} ) Am²kg⁻¹</th>
<th>( J_{is}/J_s )</th>
<th>( H_{cr} ), mT</th>
<th>( H_{cr} ), mT</th>
<th>( H_{cr}/H_c^0 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAR517e</td>
<td>1 hr</td>
<td>22</td>
<td>70</td>
<td>0.314</td>
<td>5.3</td>
<td>10</td>
<td>1.89</td>
</tr>
<tr>
<td>CAR52d</td>
<td>2 hr 16 min</td>
<td>37</td>
<td>121</td>
<td>0.306</td>
<td>5.0</td>
<td>8.8</td>
<td>1.76</td>
</tr>
<tr>
<td>CAR58f</td>
<td>3 hr 19 min</td>
<td>43</td>
<td>124</td>
<td>0.347</td>
<td>7.4</td>
<td>10.6</td>
<td>1.43</td>
</tr>
<tr>
<td>CAR519e</td>
<td>4 hr</td>
<td>55</td>
<td>130</td>
<td>0.423</td>
<td>7.7</td>
<td>13</td>
<td>1.69</td>
</tr>
<tr>
<td>CAR58d</td>
<td>4 hr 50 min</td>
<td>69</td>
<td>168</td>
<td>0.411</td>
<td>8.5</td>
<td>12.5</td>
<td>1.47</td>
</tr>
<tr>
<td>CAR519f</td>
<td>7 hr</td>
<td>61</td>
<td>142</td>
<td>0.430</td>
<td>7.9</td>
<td>13.1</td>
<td>1.66</td>
</tr>
<tr>
<td>CAR511d</td>
<td>7 hr 01 min</td>
<td>73</td>
<td>178</td>
<td>0.410</td>
<td>8.4</td>
<td>13.2</td>
<td>1.57</td>
</tr>
<tr>
<td>CAR521e</td>
<td>10 hr</td>
<td>87</td>
<td>199</td>
<td>0.439</td>
<td>8.8</td>
<td>15.4</td>
<td>1.75</td>
</tr>
<tr>
<td>CAR512d</td>
<td>10 hr 18 min</td>
<td>106</td>
<td>246</td>
<td>0.431</td>
<td>9.6</td>
<td>14.0</td>
<td>1.46</td>
</tr>
<tr>
<td>CAR512d (repeated)</td>
<td>10 hr 18 min</td>
<td>104</td>
<td>263</td>
<td>0.395</td>
<td>9.9</td>
<td>14.0</td>
<td>1.41</td>
</tr>
<tr>
<td>CAR55d</td>
<td>15 hr 01 min</td>
<td>90</td>
<td>218</td>
<td>0.413</td>
<td>8.7</td>
<td>14.7</td>
<td>1.69</td>
</tr>
<tr>
<td>CAR56e</td>
<td>21 hr 55 min</td>
<td>107</td>
<td>252</td>
<td>0.425</td>
<td>9.6</td>
<td>14.0</td>
<td>1.46</td>
</tr>
<tr>
<td>CAR57b</td>
<td>32 hr</td>
<td>117</td>
<td>299</td>
<td>0.391</td>
<td>9.4</td>
<td>15.0</td>
<td>1.60</td>
</tr>
</tbody>
</table>

Table 2. Hysteresis cycle characteristics (measured at 400°C) of 13 distinct samples, each of them having been heated previously at 400°C for a different duration (second column). The maximum applied field during hysteresis experiments was 0.190 mT. The experimental procedure is described in Section 2.2.
Figure 13. Changes in the main characteristic quantities of hysteresis cycle at 400°C versus the duration of heating time at this temperature, as measured on 13 distinct samples from hand sample CAR5. The relative magnetic homogeneity of that set of samples, attested by the regularity of the observed changes with heating time, results from the rigorous selection criteria of our samples (Section 2.1).

decrease with time, especially during the latest stage of CRM acquisition. However, the observed trend in coercivity described in Section 4.1 does not corroborate that expectation: coercive forces remain constant during that stage (Fig. 13 and Table 2). Thus, our experimental data are at variance with some theoretical inferences based on numerical micromagnetics.

The present state-of-the-art does not allow investigating the origin of this discrepancy. On one side, there are the shortcomings (like neglecting the effects of defects) and all the unsolved difficulties of micromagnetics computations which makes ‘any comparison with experiments of the computational results for perfect particles . . . meaningless and misleading’ (Aharoni 2000). On the other side, the quantitative constraints of the exsolution-and-growth mineralogical model of oxyexsolution remain to be investigated, in particular the dynamics of the propagation of oxyexsolution in the sample and individual titanomagnetite crystals, and the evolution of the shape of magnetite intergrowths as this transformation proceeds.

4.3 Blocking of chemical and thermochemical remanences

All previous theoretical studies on CRM acquisition dealt with grain-growth CRM (Stacey & Banerjee 1974; Shcherbakov & Sycheva 1996; McClelland 1996; Stokking & Tauxe 1990). The present case is more complex: both mineralogical models of exsolution (Fig. 14) indicate that both chemical composition and volume of magnetic particles do change with time, which is confirmed by the hysteresis measurements reported above (Section 4.1). Moreover the increase in volume is strictly limited by the ilmenite lamellae framework.

As pointed out by Néel (1949), regardless of the physical quantity acting (field, temperature, time etc . . .), magnetic blocking can be
Chemical and thermal remanences

Figure 14. Oxyexsolution in titanomagnetite: a comparison of spinodal decomposition and nucleation and growth (modified from Putnis 1992).

described as resulting from a single process, the increase with time of the relaxation time \( \tau \) given by:

\[
\frac{1}{\tau} = C \exp \left( -v \frac{Hc \, Is}{2kT} \right),
\]

where the coefficient \( C \approx 10^{10} \, \text{s}^{-1} \) (Aharoni 1992) estimates the frequency of ‘attempts’ to overcome the potential barrier that prevents reversal of the moment. In the present case, magnetization blocking is governed by the way titanium progressively migrates from the spinel towards and into the hexagonal corundum structure. Whichever way exsolution develops (Fig. 14), two magnetic blocking processes are at work simultaneously: (i) a progressive increase of the volume of titanium-poor titanomagnetite with a given chemical composition and (ii) a progressive increase in Fe/Ti ratio within a given volume. The increase in Fe/Ti ratio raises spontaneous magnetization, which in turn increases the microscopic coercive force in proportion. Thus, chemical change and volume increase work together towards an increase of the relaxation time.

Let us examine the following two limiting cases of CRM blocking: blocking by chemical changes with an unchanged volume of magnetic mineral (\( Is \)-blocking model) or volume-growth blocking with an unchanged chemical composition (\( v \)-blocking model). In the \( Is \)-blocking model, \( v = 2 \times 10^{-3} \, \mu \text{m}^3 \) is constant, and blocking at 400°C occurs when spontaneous magnetization reaches the value \( Is(b) = 23 \, \text{kA m}^{-1} \). This is compatible with a titanomagnetite having an ulvospinel molecular fraction \( x \) close to 0.26, a Curie temperature near 415°C, and \( Is_0 = 306 \, \text{kA m}^{-1} \) (Kawai 1956). In the \( v \)-blocking model, \( Is = 257 \, \text{kA m}^{-1} \) is constant, and the blocking volume at 400°C is \( v(b) = 0.2 \times 10^{-3} \, \mu \text{m}^3 \). For a dimension ratio equal to 2, this volume corresponds to a length of about 90 nm. Considering the development with time of titanium concentration (Fig. 14), the spinodal decomposition model tends to favour the first magnetic blocking process, while the second magnetic blocking process is more compatible with the nucleation and growth model.

In the case of TCRM blocking, all variable terms in equation 4 vary simultaneously, each of them acting as to increase relaxation time. Thus, it is difficult to separate the role of each process (chemical change, volume increase and temperature decrease) with regard to magnetic blocking. However, some constraints can be derived from thermodynamic and magnetic considerations. As pointed out above, the chemical reaction rate dependence on T is so strong that most of changes clearly occur within the highest temperature range of the TCRM experiments. Moreover, the per descendum temperature trend during blocking requires the blocked magnetic volumes to correspond, at any temperature, to the highest Curie temperatures then present in the sample. The blocking temperature spectrum (at 400°C) of the TCRM acquired by sample CAR56b has been calculated from the TCRM acquisition curve (Fig. 8b) after correction by the thermal variation of \( Js \) (measured on the adjacent sample CAR57b previously heated for 32 hr at 400°C). The TCRM blocking curve so obtained is shown on Fig. 15. This curve is somewhat approximate because of the various sources of imprecision in the \( Js(T) \) correction. In particular, the progressive slight decrease in TCRM that seems to occur from 450°C to 400°C is believed to

© 2006 The Authors, GJI, 166, 12–32
Journal compilation © 2006 RAS
There is no noticeable contribution from intermediate Curie temperature phases to the CRM. Consequently, for the subsequent Thellier experiments on CRM, the first temperature step chosen was 490°C. At the same time, beyond this temperature, each step was reduced to 4°C in order to increase the number of data in the unblocking interval of CRM. Fig. 16(b) shows an example of the unexpected behaviour then observed in the range 490–500°C. As the heating temperature is stepwise increased, a steep increase in CRM and a somewhat symmetric decrease in TRM occur. From 490°C to 502°C, the TRM is successively reduced to 66, 58, 28 and 12 per cent, that in spite of the increase of its blocking temperature range. To avoid any interference with thermal blocking/unblocking processes, another experiment was carried out, the sample being repeatedly heated at the same temperature (490°C). As seen in Fig. 17(a), the TRM(490°C, 400°C) is initially different from zero and vanishes progressively as the number n of heating increases. In contrast, the CRM(490°C, Tc) tends to increases with n. As can be seen in Fig. 17(b), the decrease of the acquired pTRM varies in proportion to the increase of the CRM left.

A possible first-order explanation of these observations is an increase in the volume of individual magnetic particles during Thellier experiments. This growth might be a mere continuation of the similar trend documented during the latest stage of CRM acquisition (Section 4.1). According to eq. 4, the expression of the blocking temperature is:

$$Tb = \frac{Hc Is}{2k \ln(C \tau_{bi})},$$

where $\tau_{bi}$ is the blocking time, which depends on cooling rate. This equation shows that a very small volume increase results in a noticeable shift of $T_b$ towards higher values. At the temperatures under consideration, a 1 per cent increase in $v$ would increase $T_b$ by 8°C. From a theoretical standpoint, an increase in volume of individual SD particles results in a similar increase in remanence intensity, regardless of the kind of remanence considered (CRM, TCRM or TRM). Since CRM and TRM have the same direction and polarity in our experiments, the CRM increase may be explained by the overstepping of the 490°C boundary by particles formerly acting as TRM carriers. This shift should also decrease the intensity of the low temperature pTRM, providing that the source of blocking temperatures less than 490°C is depleted.

As seen above, the occurrence of chemical changes within the TRM blocking range (typically 510–540°C for CRM and 470–540°C for TCRM) is indicated -both in some of the CRM-carrying samples and in the three TCRM-carrying samples- by a systematic increase in pTRM capacity (Figs 9 and 10). This trend may also be interpreted as due to an increase in particle volume. At each temperature step of Thellier experiments, chemical changes take place essentially during the first heating cycle (field off). If the volume, and thus the magnetic moment of each magnetic grain increases, so do both the pre-existing and newly acquired remanences in proportion, as long as the single domain structure is maintained. Thus, even though sample magnetic mineralogy does evolve from one temperature step to the next, it is conceivable that the TCRM/CRM ratio might not be noticeably modified. Actually, the chemical changes occurring during Thellier experiments do not seem to alter significantly the CRM/TRM ratio. For $T_r = 400°C$, this ratio is found to be $0.36 \pm 0.05$ for the two samples for which no noticeable pTRM increase is observed during Thellier experiments. It is equal to $0.36 \pm 0.08$ for the four other.

5 CHANGES DURING THELLIER EXPERIMENTS

Under laboratory conditions intended to simulate natural deuteric oxidation, titanomagnetite is not able to reach equilibrium with the oxidizing environment within a reasonable time scale, unless oxidation, titanomagnetite is not able to reach equilibrium with the high temperature treatment. That this phase starts forming rapidly, within 3 hr from the onset of heating from 400°C using small constant $\Delta T$ intervals, no significant variation in CRM intensity is generally observed until CRM starts to unblock around 490–500°C (Fig. 16a). This suggests that there is no noticeable contribution from intermediate Curie temperature phases to the CRM. Consequently, for the subsequent Thellier experiments on CRM, the first temperature step chosen was 490°C. At the same time, beyond this temperature, each step was reduced to 4°C in order to increase the number of data in the unblocking interval of CRM. Fig. 16(b) shows an example of the unexpected behaviour then observed in the range 490–500°C. As the heating temperature is stepwise increased, a steep increase in CRM and a somewhat symmetric decrease in TRM occur. From 490°C to 502°C, the TRM is successively reduced to 66, 58, 28 and 12 per cent, that in spite of the increase of its blocking temperature range. To avoid any interference with thermal blocking/unblocking processes, another experiment was carried out, the sample being repeatedly heated at the same temperature (490°C). As seen in Fig. 17(a), the TRM(490°C, 400°C) is initially different from zero and vanishes progressively as the number n of heating increases. In contrast, the CRM(490°C, Tc) tends to increases with n. As can be seen in Fig. 17(b), the decrease of the acquired pTRM varies in proportion to the increase of the CRM left.

A possible first-order explanation of these observations is an increase in the volume of individual magnetic particles during Thellier experiments. This growth might be a mere continuation of the similar trend documented during the latest stage of CRM acquisition (Section 4.1). According to eq. 4, the expression of the blocking temperature is:

$$Tb = \frac{Hc Is}{2k \ln(C \tau_{bi})},$$

where $\tau_{bi}$ is the blocking time, which depends on cooling rate. This equation shows that a very small volume increase results in a noticeable shift of $T_b$ towards higher values. At the temperatures under consideration, a 1 per cent increase in $v$ would increase $T_b$ by 8°C. From a theoretical standpoint, an increase in volume of individual SD particles results in a similar increase in remanence intensity, regardless of the kind of remanence considered (CRM, TCRM or TRM). Since CRM and TRM have the same direction and polarity in our experiments, the CRM increase may be explained by the overstepping of the 490°C boundary by particles formerly acting as TRM carriers. This shift should also decrease the intensity of the low temperature pTRM, providing that the source of blocking temperatures less than 490°C is depleted.

As seen above, the occurrence of chemical changes within the TRM blocking range (typically 510–540°C for CRM and 470–540°C for TCRM) is indicated -both in some of the CRM-carrying samples and in the three TCRM-carrying samples- by a systematic increase in pTRM capacity (Figs 9 and 10). This trend may also be interpreted as due to an increase in particle volume. At each temperature step of Thellier experiments, chemical changes take place essentially during the first heating cycle (field off). If the volume, and thus the magnetic moment of each magnetic grain increases, so do both the pre-existing and newly acquired remanences in proportion, as long as the single domain structure is maintained. Thus, even though sample magnetic mineralogy does evolve from one temperature step to the next, it is conceivable that the TCRM/CRM ratio might not be noticeably modified. Actually, the chemical changes occurring during Thellier experiments do not seem to alter significantly the CRM/TRM ratio. For $T_r = 400°C$, this ratio is found to be $0.36 \pm 0.05$ for the two samples for which no noticeable pTRM increase is observed during Thellier experiments. It is equal to $0.36 \pm 0.08$ for the four other.

Figure 15. TCRM blocking (dashed line, calculated), TCRM unblocking (solid line, measured), and TRM blocking (solid line, measured) for sample CAR56b.

be an artefact. However, the interval of TCRM blocking is clearly established: more than 90 per cent of remanence is blocked between 540°C and 500°C. This indicates that TCRM is carried by phase 1 mineral. The magnetic measurements represented on Fig. 8(b) show that this phase starts forming rapidly, within 3 hr from the onset of the high temperature treatment.
6 COMPARING CHEMICAL AND THERMAL REMANENCES: FROM THEORY TO PRACTICE

6.1 Theoretical considerations

According to Néel (1949), the average magnetic moment of a set of identical non-interacting SD particles with their easy magnetization axis parallel to the applied field $H$, when measured at any temperature $T_0$ equal or less than $T_b$, is:

$$m_0 = v I_s J_0 \frac{v I_s H}{k T}.$$  \hfill (6)

For a randomly oriented assembly integrated from 0 to $\pm \pi/2$, and considering that the argument of the hyperbolic tangent is less than 1 in our case, the total moment at $T_0$ is approximately given by:

$$M_0 = \frac{1}{3} v I_s J_0 \frac{v I_s H}{k T}.$$  \hfill (7)

Applying the blocking equation (4), we can replace $T = T_b$ by $v Hc I_s/2k \ln(C \tau(b))$ where $\tau(b)$ is the blocking time of the remanent magnetization under consideration. Also, since the coercive force is due to shape anisotropy in the present case, and provided magnetization reversals are coherent, $Hc$ is equal to $Nd I_s$. Then the remanence intensity is expressed by:

$$M_r = \frac{2}{3} H \tau(b) \frac{\ln(C \tau(b))}{I_s}.$$  \hfill (8)

Figure 16. CRM and TRM changes versus temperature during Thellier experiments in case of (a) 10°C step heating from 400°C to 550°C or (b) a rapid heating (4°C/min) to 492°C followed by 4°C step heating up to 546°C.
Figure 17. Effect of repetitive double heatings at 490°C on the CRM(490°-Tc) and the TRM (490°–400°) of sample CAR518c (a). Plot of TRM (490°–400°) versus CRM(490°-Tc) (b). The dot labels indicate the sequence number of successive double heatings.

For CRM, the expression may be written:

\[ M_{CRM} = \frac{2 H v I_s}{3 N d} \ln \left( C \tau_{CRM} \right) \frac{I_s(CRM)}{I_s}, \]

where, as usual (Néel 1949; Aharoni 1992), the blocking time \( \tau_{CRM} \) is taken to be of the same order as the time taken for the measurement (14 s in the present case). In eq. (9) all the quantities are initially associated with the magnetic carrier at the blocking instant. At this moment, \( v \) and \( I_s \) are largely different for the two blocking models considered above (Section 5), and so is the CRM intensity. Subsequently, however, since the mineralogical process is still going on and is evolving towards the same final mineral assemblage, CRM intensity converges towards the same value for the volume blocking and the \( I_s \)-blocking models.

If a TRM is now imparted to this final assemblage, and assuming that no changes occur during TRM acquisition, we have similarly:

\[ M_{TRM} = \frac{2 H v I_s}{3 N d} \ln \left( C \tau_{TRM} \right) \frac{I_s(CRM)}{I_s}. \]

where \( I_s(CRM) \) is the spontaneous magnetization of the final mineral produced during CRM experiments, measured at \( T_{CRM} \), the temperature of TRM blocking. \( \tau_{TRM} \), the blocking time for thermoremanence, is the time needed for \( \tau \) to increase sufficiently for a superparamagnetic grain population initially close to the SP/SD threshold to become fully blocked (Néel 1949; Dodson & McClelland-Brown 1980). Thus \( \tau_{TRM} \) is inversely proportional to cooling rate. This is the origin of the dependence of TRM intensity upon cooling rate, first experimentally verified by Papusoi (1972).
It follows from eqs (9) and (10) that:

\[
\frac{M_{\text{crm}}}{M_{\text{trm}}} = \frac{I_s(\text{Trm})}{I_s(\text{Term})} \ln(\frac{C(\text{crm})}{C(\text{trm})}).
\]

(11)

This equation is equivalent to the expressions obtained by Stacey & Banerjee (1974), Stokking & Tauxe (1990) and Shcherbakov & Sycheva (1996) who all neglected the logarithmic term.

For the TCRM case, we have similarly:

\[
\frac{M_{\text{crm}}}{M_{\text{trm}}} = \frac{I_s(\text{Trm})}{I_s(\text{Term})} \ln(\frac{C(\text{crm})}{C(\text{trm})}).
\]

(12)

where \( \tau_{(\text{term})} \) is the blocking time for TCRM. In eqs (11) and (12) expressions such as \( I_s(\text{Term}) \) has the meaning of ‘spontaneous magnetization at the TRM blocking temperature’.

It must be reminded that eqs (8)–(10) are correct only if the mode of magnetization reversal of particles is coherent. If not, the magnetic moment calculated from these expressions should be too low. However, as discussed in Section 4.2, some of our experimental evidences do not seem to support the occurrence of non-coherent reversal in the CRM-carrying particles. In any case, the occurrence of incoherent magnetization reversals would not affect the validity of eqs (11) and (12) since they express the ratio of two magnetic moments.

6.2 Comparing experimental data and theoretical predictions

The experimental TCRM/TRM average is 1.11 ± 0.05. According to the VTM measurements carried out on sample CAR56b (Fig. 15), the median blocking temperatures of TCRM and TRM are very close to each other. Considering the uncertainties in the method of calculation of the TCRM blocking curve (Section 4.3) and the fact that repetitive heating does not seem to modify the Curie temperature of phase 1 mineral (Fig. 12), the TCRM- and TRM blocking temperatures may be assumed to be equal. Regarding the blocking time \( \tau \), Néel (1949) proposed to define it as the time needed for the relaxation time to be multiplied by \( e \). Thus, in our case \( \tau_{(\text{term})} \) is inversely proportional to both the cooling rate and the rate of \( I_s \) increase and volume growth. Since the rates of the physical and chemical changes of grains are unknown, a maximum estimate of \( \tau_{(\text{term})} \) is used, based on the cooling rate alone (0.1°C/min). The TRM cooling rate is 4°C/min. Using those values of \( T_b(\text{term}) \) and \( \tau \), one obtains a theoretical TCRM/TRM ratio equal to 1.1, which is in agreement with experimental findings. However, this perfect coincidence is probably fortuitous: any small difference between \( T_b(\text{term}) \) or \( T_b(\text{term}) \) would result in largely different theoretical TCRM/TRM ratios.

For samples carrying a CRM, the experimental CRM/TRM ratios is found to increase with temperature \( T_c \) (0.36 ± 0.07 at 400°C, 0.66 ± 0.02 at 450°C, and 0.90 ± 0.02 at 500°C), which is in agreement with the trend expected from eq. (11). These experimental findings seem to be of general validity. Nguyen & Pecherskiy (1985) reported a CRM/TRM ratio equal to 0.47 for a CRM acquired by apparently single domain magnetite resulting from oxysolution of large-sized basaltic titanomagnetite heated in air for 26 hr at 400°C. In the absence of Thellier experiments, Nguyen & Pecherskiy (1985) calculated the CRM/TRM ratio from the values of initial CRM and final TRM. According to our Thellier experiments, this method provides CRM/TRM ratios larger (by 20–25 per cent) than those obtained from the linear part of CRM–TRM diagrams (Fig. 9). Thus their experimental data and ours are in good agreement. We can note in passing that there is no reason for the size of the exsolved phases to be dependent on the size of the host titanomagnetite. Rather, it should be generally expected that exsolution size and spacing are mainly dependent on the temperature of oxysolution because of the strong dependence of the rate of ion diffusion on temperature.

The theoretical CRM/TRM values were calculated using a median TRM blocking temperature of 525 ± 5°C, as typically found from our Thellier experiments. As can be seen on Fig. 11, theoretical ratios are systematically and significantly lower than experimental determinations: 0.12 ± 0.04 at 400°C, 0.17 ± 0.05 at 450°C, and 0.39 ± 0.11 at 500°C.

For CRM, another inference from the theory of magnetism of non-interacting single domain is not experimentally verified. The very same eq. (11) that qualitatively accounts for the observed trend in CRM/TRM ratio as the blocking temperature of CRM is modified, also implies that the Thellier experiments provide concave up CRM–TRM plots. The concavity expected from theory is large enough to be easily seen in Fig. 9. For a CRM acquired at 400°C, the slope of the curve in the TRM blocking range should be equal to 0.26 near the lower limit (typically 510°C) and while it should be 0.02 near the upper limit (typically 540°C). Actually, all our samples exhibit a nice linear plot over almost the entire TRM blocking range, whether pTRM checks are positive or not.

6.3 Possible effects of magnetic interactions

The differential hysteresis experiments carried out at room temperature, especially the FORC diagram, demonstrate the presence of large magnetic interactions in between the single-domain CRM-carrying particles. This conclusion certainly holds for TCRM-carrying particles as well. It is well established that magnetic interactions can largely reduce the intensity of low-field remanences such as TRM or ARM (Dunlop & West 1969), and a similar effect has to be expected for CRM. In the present context, which we wish to know is whether magnetic interactions modify the ratio of CRM to TRM. Shcherbakov et al. (1996) addressed this topic by carrying out a comparative Monte Carlo simulation of CRM and TRM acquisition by SD interacting uniaxial magnetite grains. For a crystallization temperature of 300°C, they found that CRM/TRM ratios less than 1 were typical of non-interacting grains with volume concentration less than 1 per cent. For concentrations larger than 1–3 per cent, CRM/TRM ratios equal to or larger than 1 were computed. In our basaltic samples, the almost-magnetite formed by oxysolution (our phase 1) is clustered within the former Ti-rich titanomagnetite crystals where its concentration is very large. Considering that these initial Fe-Ti-O crystals contained some 70–80 per cent of ulvospinel before oxidation (Section 3.1), the local volumetric concentration of magnetite should lie in the range 20–30 per cent. According to the theoretical results of Shcherbakov et al. (1996), a CRM/TRM ratio close to 1 or more has therefore to be expected for our samples, with consequently no dependence with the temperature of CRM acquisition. These predictions are at odds with our experimental findings and those of Nguyen & Pecherskiy (1985) as well.

However, the simple qualitative considerations developed below suggest that magnetic interactions can explain the two major discrepancies noted above between experimental data and predictions from the non-interacting single-domain theory of magnetism: (i) experimental CRM/TRM ratios systematically larger than theoretical ones and (ii) experimental CRM–TRM diagrams linear (Fig. 9) rather than concave up.
Whichever mineralogical scenario of oxyexsolution is considered (Fig. 14), weak magnetic interactions are expected at the instant when the direction of the CRM-carrying SD moment gets blocked. In case of spinodal decomposition, the remanence-carrying particles have then a very weak spontaneous magnetization (Section 4.3). In case of nucleation and growth, the blocked particles are strongly magnetic, but the intragranular distance is generally much larger than the grain diameter. Subsequently, spontaneous magnetization or particle size increases so that interaction field grows up, reaching 45 mT after completion of thermal treatments (Section 4.2). These interactions are about three times weaker than particle axial coercivity and cannot significantly reduce the intensity of the previously blocked CRM. However, they should reduce the intensity of the TRM initiated later in this highly interactive magnetic configuration.

The linearity of experimental CRM–TRM diagrams can also result from magnetic interaction. The increase of these fields as oxyexsolution proceeds should result in a random redistribution of the intrinsic blocking temperatures of grains. This suggestion is in agreement with Néel’s (1949) theory of magnetism of SD particles which implies that the energy barrier $E$ of a particle depends on external fields. $E$ increases when the external field $H$ is parallel to the magnetic moment of particle and decreases when they are antiparallel, the amplitude of the change in $E$ increasing with the magnitude of $H$. This results in qualitatively similar shifts of $T_b$. When subjected to strong interaction fields of different sign and magnitude, a set of identical single domain grains with an initially unique intrinsic $T_b$ yields a broad distribution of $T_b$’s around the initial value. According to our experiments, interaction fields appear large enough for redistributing $T_b$’s all over their intrinsic (interaction-free) range of variation.

Regarding TCRM, the agreement between experimental and theoretical remanence ratios suggests that large interaction fields are already established before TCRM blocking. This suggestion implies that oxyexsolution develops largely before blocking, that is to say during the 3-hr cooling from 560°C to the upper limit of the TCRM blocking range, close to 540°C (Fig. 15). This view is supported by the fact that at a temperature as low as 400°C, the breaking down of titanomaghemite into a magnetite/ilmenite assemblage does not take more than 3 hr, while theoretical considerations suggest that the reaction rate in the 540°–560°C range is larger than that at 400°C by 2 to 3 orders of magnitude (Section 4.1). It is thus reasonable to assume that most of mineralogical changes do occur before TCRM blocking. The behaviour of TCRM should then tends towards that of a simple TRM, in agreement with the fact that the TCRM/TRM ratio is found to be equal to 1 (once the cooling rate difference is taken into account) and the TCRM versus TRM plot is linear.

### 7 CONCLUSIONS

The present study suggests that isothermal oxyexsolution process in basaltic titanomagnetite at moderate temperatures (500°–400°C) results from nucleation and growth rather than from spinodal decomposition. Tiny Ti-poor titanomagnetite domains of nanometric size form rapidly along the boundaries of nucleating ilmenite lamellae. The magnetization of these domains gets blocked in the absence of noticeable magnetic interactions, due to the large distance between them. The orientation of magnetic moments remains constant while the size of the single domain Ti-poor titanomagnetite develops within the ilmenite lamellae framework and interaction fields strongly increase. The final product is an assemblage of strongly coercive (110–140 mT) single-domain particles with strong magnetic interactions between them (around 45 mT). As a result of these local interaction fields, the intrinsic blocking/unblocking temperatures of individual particles are modified in a random fashion. In our experiments, this effect results in linear CRM/TRM curves, in contradiction with the concave up shape expected from the theory of magnetism for non-interacting single-domain grains. The alteration of Tb distribution during oxyexsolution help to explain Walderhaug’s (1992) observation that the genuine direction of the primary remanence of rocks heated in a weak field for several hours at moderate temperatures (400°–525°C) can no longer be retrieved after CRM acquisition. This overlap of the blocking temperatures can result from the redistribution of the blocking temperatures of the primary remanence under the influence of stronger and stronger interaction fields. In nature, long-term heating at moderate temperatures are expected to occur typically near igneous contacts. It is often observed that primary and secondary magnetizations cannot be separated from each other in the baked zone, as pointed out by Camps et al. (1995) for the Steens Basalt sequence. This suggests that such magnetic overprints are low-temperature chemical remanences rather than partial thermoremanences.

In agreement with previous experimental works (Nguyen & Pecherskiy 1985; Stokking & Tauxe 1990), our experiments show that the intensity of the isothermal CRM acquired at a moderate temperature $T_r$ by single domain particles is largely inferior to that of the total TRM acquired above this temperature. The CRM/TRM ratio is shown here to be strongly dependent on temperature $T_r$, with values from less than 0.4 at 400°C to 0.9 at 500°C. These findings are in qualitative agreement with the theory of non-interacting single domain particles. However, the experimental ratios are significantly larger than theoretical values, which we tentatively attribute to the effect of magnetic interactions.

Knowing that CRM can provide a linear CRM–TRM plot when studied by the Thellier method invites particular caution in paleointensity studies on geological materials which may have experienced late mineral crystallization at moderate or low temperatures, as for example baked contacts or volcanic glasses. Because they generally provide nice linear NRM–TRM diagrams, baked contacts are often considered as an ideal material for paleointensity studies. However, considering how moderate is the temperature reached during base- ment heating by a cooling lava or dyke (Audunsson & Levi 1988; Camps et al. 1995; Kristjansson 1985) and the long duration of this process, there are good reasons to have some doubts about the actual nature of NRM. Volcanic glasses constitute another material of debatable reliability for paleointensity studies. Prone to rapid alteration at water temperature (Jakobsson 1978; Moore 1966), they are also thermodynamically unstable and can undergo post-eruption devitrification (e.g. Marshall 1961). Submarine basalt glasses (SBG) generally provide linear NRM–TRM plots (Pick & Tauxe 1993) and are thus particularly appreciated for paleointensity studies. However, it is puzzling that the remanence of SBG is carried by some almost stoichiometric magnetite (Pick & Tauxe 1994) instead of high-titanium titanomagnetite, the pre-deuteric magnetic phase system-atically present in fresh submarine pillow basalts (Schaeff er & Schwarz 1970; Prévot et al. 1979) and continental tachylites as well (Smith & Prévot 1977). It is well established (e.g. O’Reilly 1984) that Ti-poor titanomagnetite can crystallize at seawater temperature and thus carry a CRM rather than a TRM. In contrast, basaltic high-titanium titanomagnetite have crystallization temperature largely exceeding their Curie temperature.

Heller et al. (2002) showed that the Earth’s mean dipole moment calculated from the paleointensity data obtained from SBG over the
last 20 Ma is unexpectedly 40 per cent less than the moment calculated from the data provided by continental volcanic rocks, although the main magnetic carriers are typically Ti-poor titanomagnetite in both cases. Thus, these authors suggested that a significant fraction of SBG provides too low palaeointensity determination due to a low-temperature crystallization of magnetic carrier. The results described in the present paper give a solid experimental support to their suggestion.

Most of palaeointensity data from continental volcanic rocks are obtained from basaltic flows containing titanomagnetite oxysexolved as a result of deuteric alteration during magma cooling. The TCRM experiments reported in the present paper are the first attempt to mimic this natural phenomenon in order to evaluate its possible effects on palaeointensity determination. Our experimental data suggest that titanomagnetite oxysexolution concomitant with magnetization blocking results in TCRM/TRM ratios larger than 1 by some 10 per cent. Theoretical considerations suggest that this overestimate might be simply explained by the difference in cooling rates during TCRM and TRM acquisition processes in the laboratory. In nature, it is probable that oxysexolution develops mostly at temperatures higher than the TRM blocking range. In this case, no bias due to magnetic interaction should affect palaeointensity data. However, the difference in cooling rate between the natural and laboratory blocking processes has to be taken into account according to expression 12.

ACKNOWLEDGMENTS

We thank our colleagues at the Saint Maur geomagnetic laboratory, especially Mansour Bina, and at the Institute for Rock Magnetism (IRM) in Minneapolis, Minnesota, for helping UD during her stays in these laboratories. The choice of the rock used here was based on low-field thermomagnetic studies previously carried out by Elisabeth Schneppe, who kindly provided some of her material to us. We acknowledge Pierre Rochette and Fabienne Vadeboin at CEREGE (University Aix-Marseille and CNRS) who kindly carried out FORC and differential IRM experiments and computation, and Liliane Faynot for her help in drawing figures. Ken Hoffman kindly revised an early version of this paper and made both linguistic and scientific suggestions for improvement. Reviews by David Dunlop, Ron Merrill and an anonymous referee led to substantial and scientific suggestions for improvement. Reviews by David

REFERENCES

Dunlop, D.J., 1987. Temperature dependence of hysteresis in 0.04–0.22 μm magnetite and implications for domain structure, Phys. Earth planet. Inter., 46, 100–119.