

A new chlorite geothermometer for diagenetic to low-grade metamorphic conditions

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1	Revision 1
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26	
27	Abstract
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29	The evolution of chlorite composition with temperature (and pressure) serves as basis to a
30	number of chlorite chemical thermometers, for which the oxidation state of iron has been
31	recognised as a recurrent issue, especially at low temperature (T). A new chlorite
32	geothermometer that does not require prior Fe ³⁺ knowledge is formulated, calibrated on 161
33	analyses with well constrained T data covering a wide range of geological contexts, and tested
34	here for low- T chlorites ($T < 350$ °C and pressures below 4 kbar). The new solid-solution
35	model used involves six end-member components (the Mg and Fe end-members of 'Al-free

36 chlorite S', sudoite and amesite) and so accounts for all low-*T* chlorite compositions; ideal
37 mixing on site is assumed, with an ordered cationic distribution in tetrahedral and octahedral

38 sites.

39 Applied to chlorite analyses from three distinct low-T environments for which independent T

data are available (Gulf Coast, Texas; Saint Martin, Lesser Antilles; Toyoha, Hokkaido), the

new pure-Fe²⁺ thermometer performs at least as well as the recent models, which require an

estimate of Fe³⁺ content. This relief from the ferric iron issue, combined with the simple

formulation of the semi-empirical approach, makes the present thermometer a very practical

tool, well suited for, e.g., the handling of large analytical datasets – provided it is used in the

calibration range (T < 350°C, P < 4 kbar).

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Keywords: Chlorite, Geothermometry, Diagenesis, Low-grade metamorphism, Solid-solution.

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Introduction

- 51 Chlorites are ubiquitous in most diagenetic and metamorphic rocks (Foster 1962;
- 52 Cathelineau 1988; Laird 1988; De Caritat et al. 1993; Vidal et al. 2001). Their wide
- compositional variations, through the effect of the three main substitutions $Fe^{2+} = Mg$,
- Tschermak $Al^{IV}Al^{VI} = Si(Mg, Fe^{2+})$, and di/trioctahedral $3(Mg, Fe^{2+}) = \Box + 2Al^{VI}$ (where \Box
- represents an octahedral vacancy), is known to be sensitive to the pressure (P) and
- 56 temperature (*T*) conditions of formation, to the bulk composition and to the physico-chemical
- properties of the environment, like activities in the fluid(s) (e.g. Cathelineau and Nieva 1985;
- Hillier and Velde 1991; Vidal and Parra 2000). This makes chlorites good indicators of rock
- 59 history and, for this reason, they are the basis of several empirical or thermodynamic methods
- to estimate *P-T* conditions (e.g. Cathelineau 1988; Hillier and Velde 1991; Walshe 1986;
- 61 Vidal et al. 2001; Inoue et al. 2009).
- Several studies showed however that an empirical approach, in which the temperature
- is directly linked to chlorite composition, is often inappropriate (e.g. De Caritat et al. 1993;
- Essene and Peacor 1995; Bourdelle et al. *in revision*). For example, the empirical calibrations
- 65 that are based on the T-dependent variation of Al^{IV} content are easily applicable, but do not
- account for the influence of bulk-rock composition. Thus, this approach has no generality and
- can only be applied to a restricted range of geological settings (De Caritat et al. 1993), in spite
- of the many formulations proposed (Cathelineau and Nieva 1985; Kranidiotis and McLean
- 69 1987; Cathelineau 1988; Jowett 1991; Hillier and Velde 1991; Zang and Fyfe 1995; Xie et al.
- 70 1997). Thermodynamic methods circumvent this problem, allowing one to account for

mineral assemblage and/or physicochemical parameters in addition to temperature and pressure (e.g. Walshe 1986; Vidal and Parra 2000; Vidal et al. 2001, 2005, 2006). However, the thermodynamic approach may be also questioned, considering that the large range of compositional variations in chlorites was not properly handled by thermodynamic modelling. In fact, the inaccuracy of this approach is due in most cases to uncertainties in the *P-T* data employed for the calibration, to the inaccuracy of the thermodynamic properties of chlorites end-members, to the choice of the mixing model, and to the difficulty in measuring the Fe³⁺ content and in accounting for it in solid-solution models (cf. Grosch et al. 2012).

On this basis, Inoue et al. (2009) proposed a new, better suited low-*T* geothermometer that does not require the knowledge of end-member thermodynamic properties but includes the effect of Fe³⁺ content on temperature estimation, thereby requiring independent Fe³⁺ determination. However, in their comparative study based on high-spatial-resolution analysis, Bourdelle et al. (*in revision*) obtained some unsatisfactory results with this thermometer for low-*T* conditions, probably due to the difficulty in obtaining reliable values of XFe³⁺ at nanometer scale. Indeed, the estimation of Fe³⁺ content needs either a Mössbauer spectroscopy study (e.g. Beaufort et al. 1992), which involves grinding of the sample and loss of the textural information, or *in situ* XANES analysis (e.g. Vidal et al. 2006) which remains a complex procedure, or a numerical estimation using an iterative multi-equilibrium calculation (Vidal et al. 2006). Thus, the presence of ferric iron is clearly an important issue.

The present study was undertaken as a continuation of previous works of Vidal et al. (2006), Inoue et al. (2009) and Bourdelle et al. (*in revision*), with the goal of developing a chlorite geothermometer for diagenetic to low-grade metamorphic conditions but avoiding the problem of estimating the oxidation state of iron in chlorite. The new calibration is based on a bibliographic compilation of data published during the past three decades and addressing a large variety of geological environments. The validity of this geothermometer is then tested by comparison of the results with those obtained with Vidal et al. (2006) and Inoue et al. (2009) thermometers on three *other* sets of analytical data, for which Fe³⁺ contents are known: chlorite analytical electron microscope (AEM) analyses from the Gulf Coast (Texas), where Fe³⁺ content was estimated by multi-equilibrium calculation; chlorite electron-microprobe (EMP) analyses from Saint Martin (Lesser Antilles), where Fe³⁺ content was estimated by Mössbauer spectroscopy (Beaufort et al. 1992); and chlorite EMP analyses from Toyoha (Hokkaido), where Fe³⁺ content was estimated by X-ray photoelectron spectroscopy (XPS, Inoue et al. 2010).

Sources of chlorite analyses

Literature data for calibration

In order to calibrate the new geothermometer on a wide spectrum in terms of P-T conditions, of geological environments and of analytical methods, published data concerning the low-T chlorites and their chemical compositions were screened. The specifications governing the selection of the data were as follows:

- A detailed quantitative analysis of chlorites by EMP or AEM should be available along with temperature and pressure data estimated by independent methods, in the range of 50-350°C and below 4 kbar. The temperature data were used as published when given with a sufficient level of confidence (i.e. convergence of several thermometric methods or discussion of the obtained data). The pressure data refer to the authors' estimates whenever possible and to indications of geological context ensuring that formation pressure was less than 4 kbar. In case of uncertainty or insufficient level of confidence regarding the *P-T* data, the analyses were discarded. For instance, some important studies such as Jowett (1991), Zang and Fyfe (1995) and Inoue et al. (2009) had to be discarded because pressure estimates were not available or because temperature data were not assigned to specific chlorite compositions. All relevant published data/studies are
- 126 -
 - as in Rahn et al. (1994) study, where authigenic and detrital compositions are identified.

 All published data summarized in Table 1 are considered as referring to authigenic chlorites.

Only analyses of material that was not identified as detrital by the authors were retained,

- chlorites.
- 130 Only quartz-bearing samples were considered.

summarized in Table 1.

When these first criteria were met, two additional chemical criteria were applied to exclude contaminated analyses or those that cannot be expressed as a linear combination of end-members. These are:

- $135 K_2O + Na_2O + CaO < 1\%wt$
- 136 Al^{VI} (+ Fe^{3+ VI}) Al^{IV} = vacancies x 2

As a result, a total of 161 analyses and relevant formation conditions were retained for the calibration (Figure 1).

Datasets for testi	ng the thermometers
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Gulf Coast chlorites

These chlorites are from sandstone core samples from nine wells of the Gulf Coast, Texas. Analyses were obtained with AEM on rock ultrathin sections obtained by focused ion beam (FIB) milling, in accordance with the analytical protocol described by Bourdelle et al. (2012). All details regarding the location and geology methods are given by Bourdelle et al. (in revision). In the following, four analyses of the crystal rim were used for each sample because these compositions were considered to be a good representation of the last steps of dissolution/recrystallisation processes and were assumed to represent the closest approach to the equilibrium composition for the highest P-T conditions. The independently obtained P-T formation data are bottom-hole temperature (BHT) and bottom-hole pressure (BHP) data corrected following Kehle (1971) and assumed to be the maximum P-T conditions (100-230°C, 0-1.2 kbar) undergone by the samples, with an estimated error range of ±20°C and ± 0.2 kbar (for discussion, see Bourdelle et al. in revision). The chemical composition of the selected Gulf Coast chlorites spans a large range (Table 2), but similar to that reported by previous studies on diagenetic clays (e.g. Curtis et al. 1984, 1985; Velde and Medhioub 1988; Hillier and Velde 1991; Jahren and Aagaard 1989, 1992; Jahren 1991). If all iron is taken as Fe²⁺, total Al contents range from 2.52 to 3.38

(Table 2), but similar to that reported by previous studies on diagenetic clays (e.g. Curtis et 1984, 1985; Velde and Medhioub 1988; Hillier and Velde 1991; Jahren and Aagaard 1989, 1992; Jahren 1991). If all iron is taken as Fe²⁺, total Al contents range from 2.52 to 3.38 atoms per formula unit (apfu) based on a 14 oxygen anhydrous basis (which will be used throughout this paper), vacancies from 0.12 to 0.49 apfu while the Fe²⁺/(Fe²⁺+Mg²⁺) ratio varies between 0.47 and 0.88 (Table 2).

The Fe³⁺ content was estimated using the approach of Vidal et al. (2006), which is based on the convergence of four chlorite-quartz-water equilibria. This method gives the minimum Fe³⁺ proportion required to reach the best convergence at specific *P-T* conditions; a maximum Fe³⁺ proportion can be estimated when the equilibrium convergence is lost. Under low-*T* conditions, the difference between minimum and maximum Fe³⁺ contents is small and, therefore, we considered that the minimum Fe³⁺ content estimated with this approach is a fair approximation of the actual value. The results show that the minimum Fe³⁺ contents are heterogeneous in the temperature window of the Gulf Coast and range between 5% and 40% of total iron.

Saint Martin chlorites

Relevant analyses have been published by Beaufort et al. (1992). The samples are from volcanoclastic rocks that were hydrothermally altered by an intrusive quartz-diorite pluton emplaced during the Early Oligocene (Beaufort et al. 1990, 1992). The chlorites used in this study are from three different alteration zones: the epidote + chlorite + quartz assemblage zone (*zone 2*), the epidote + chlorite + hematite + quartz assemblage zone (*zone 3*), and the chlorite + phengite + magnetite assemblage zone (as vein, *zone 4*). Paleotemperatures were estimated by the fluid-inclusion method, and range between 220 and 340°C (Beaufort et al. 1992; Inoue et al. 2009).

Under the assumption that $Fe_{total} = Fe^{2+}$, the Al^{IV} content ranges from 0.96 to 1.21 apfu, vacancies from 0.05 to 0.20 apfu, and the $Fe^{2+}/(Fe^{2+}+Mg^{2+})$ ratio between 0.08 and 0.68. The XFe^{3+} ratios were estimated by Mössbauer spectroscopy (Beaufort et al. 1992) and the results strongly depend on the metamorphic zones: 25-30% for zone 2 chlorites, 32% for zone 3, and 16% for zone 4.

Toyoha chlorites

The seven analyses used concerning Toyoha chlorites have been published by Inoue et al. (2010) and have been also separated from the literature data because they are associated with precise XFe³⁺ estimations. These chlorites are from hydrothermally altered rocks in the Toyoha geothermal system, southwestern Hokkaido, Japan, from the propylitic zone and the ore mineralized zone. Paleotemperatures were estimated with Inoue et al. (2009) model and were close both to the homogenisation temperatures of fluid inclusions and the present subsurface temperatures measured through drill holes. They range between 159 and 264°C.

Under the assumption that $Fe_{total} = Fe^{2+}$, the Al^{IV} content ranges from 0.85 to 1.20 apfu, Al^{VI} from 1.08 to 1.33 apfu and vacancies from 0.25 to 0.40 apfu. The Fe^{3+} contents were estimated by X-ray photoelectron spectroscopy (XPS) and the results are homogeneous and range from 0.13 to 0.17 for the ore mineralized zone and from 0.20 to 0.26 for the propylitic zone.

Formulation of a new geothermometer, as an extension of previous methods

To establish a new thermometer, it is necessary to define the cationic repartition in the structure, the activity model used and appropriate solid solutions by the choice of relevant end-member components. In this study, the focus is on chlorites formed at low-temperatures, i.e. between 50 and 350°C, which have high Si content and octahedral vacancies due to

211 $Si \square R^{2+}$ exchange (e.g. Hillier and Velde 1991). Vidal et al. (2006), building on Vidal et al. 212 (2001) and Vidal et al. (2005) studies, chose clinochlore, daphnite, Mg-sudoite, Fe-amesite 213 and Mg-amesite as end-members. This excludes Si-rich (Si > 3 apfu) compositions (Table 3) 214 and therefore, many of the diagenetic chlorites. Inoue et al. (2009) preferred to consider a 215 compositional space defined by the Mg-chlorite S, Mg-amesite, daphnite and Mg-sudoite end-216 members (Table 3). This choice seems appropriate to cover all the compositional range of 217 low-T chlorites, especially because the octahedral vacancies rarely exceed 1 apfu. However, 218 all iron is then concentrated in one trioctahedral end-member component, daphnite. 219 Considering these observations, the chlorite solid solution is defined in this study with six 220 end-members: (Fe, Mg)-chlorite S, (Fe, Mg)-sudoite and (Fe, Mg)-amesite (Table 3). 221 The activity model depends on the cationic site repartition in the structure; according 222 to the ideal structure of the trioctahedral chlorite defined by Bailey (1988), the basic layer 223 consists in a regular alternation of talc $\{M1(M2)_2[(T1)_2(T2)_2]O_{10}(OH)_2\}$ and brucite layers 224 {(M3)₂M4(OH)₆}. Each crystallographic site entails some cationic preferences, but two 225 assumptions remain possible: that of an ordered distribution (Vidal et al. 2005, 2006) and that of random mixing (Inoue et al. 2009). Vidal et al. (2006) assumed that Al^{IV} is restricted to T2 226 sites, vacancies to M1, Al^{VI} fills M4 first then M1 and eventually M2-M3, while Fe-Mg fills 227 228 M2-M3 sites then M1. Inoue et al. (2009) borrowed the distinction between T1 and T2 sites, with Al^{IV} exclusively on T2, but assumed a random mixing on all octahedral sites. The 229 230 ordered approach seems to be closest to Bailey's (1988) observations and makes Vidal's thermometer less sensitive to Fe³⁺ content, as shown in a comparative study (Bourdelle et al. 231 232 in revision). For this reason, an ordered model was adopted in this study to describe the 233 compositional variations of low-T chlorite, as follows. To account for 'chlorite S' end-234 members, we unified M1 and M4 sites (Table 4). Tetrahedral Al is restricted to T2 sites, vacancies to M1-M4, Al^{VI} resulting from Tschermak exchange fills M1-M4, excess Al^{VI} is in 235 M2-M3, Fe-Mg fills predominantly M2-M3 sites then M1-M4 (Table 4). In accordance with 236 237 Vidal et al. (2005, 2006), Fe/Mg and Si/Al ratio are considered equal in the M1-M4 and M2-M3 sites, and T2 sites respectively. Regarding the special case of Fe³⁺, Vidal et al. (2005, 238 2006) restricted it to M4 and Inoue et al. (2009) exchanged it with AlVI in all M sites (with an 239 identical Fe³⁺/Fe²⁺ ratio), whereas the new model does not consider it. 240 241 From the above cation site repartition, end-members ideal activities can be calculated 242 following the mixing-on-site model of Helgeson et al. (1978) and Powell (1978):

$$244 a_j^{ideal} = k \prod_s \prod_m (X_m)^{r_m} (1)$$

246 where a_{ideal} is the ideal part of the activity, r_m and X_m are the number and the mole fraction of 247 cation m on the site s, and k is the normalization constant defined by:

248

$$249 k = \prod_{s} \prod_{m} \left(\frac{n_s}{r_m} \right)^{r_m} (2)$$

250

- As done by Inoue et al. (2009), neglecting the non-ideal contribution of the site mixing, a new semi-empirical $T = f(\log K)$ geothermometer can be formulated with our specific choice of ordered solid-solution model, where K is the equilibrium constant of the end-member-component reaction describing the chlorite + quartz equilibrium.
- From a thermodynamic point of view, for any balanced chemical reaction involving *j* endmembers, the equilibrium condition can be expressed by:

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258
$$\Delta G_r^0 + R.T. \ln K = 0$$
 (3)

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where ΔG_r^0 is the Gibbs free energy of reaction, R is the gas constant and K is the chlorite + quartz equilibrium constant, which can be written as:

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$$263 K = \prod_{j} \left(a_{ideal} \right)_{j}^{v_{j}} (4)$$

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with v_j as the stoichiometric reaction coefficient of phase j. The equations for the calculations of end-member ideal activities are listed in Table 3, according to the cationic site mixing repartition detailed in Table 4. With three end-members, the chlorite + quartz assemblage can be described in the Mg system by the reaction:

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270 Mg-Chl S + 3 Mg-Sud =
$$3 \text{ Mg-Am} + 7 \text{ Qz} + 4 \text{ H}_2\text{O}$$
 (5)

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If the effect of pressure is ignored, in accord with the P < 4 kbar data selection, and the change in heat capacity is assumed to be naught, the log K for the reaction defined above can be expressed by:

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$$\log K = \frac{-\Delta G_r^0}{2.303(RT)} = \frac{A}{T} + B = \log \left(\frac{a_{Mg-Am}^3 a_{SiO_2}^7 a_{H_2O}^4}{a_{Mg-ChlS} a_{Mg-Sud}^3} \right)$$

$$= 3\log a_{Mg-Am} - \log a_{Mg-ChlS} - 3\log a_{Mg-Sud} + 7\log a_{SiO_2} + 4\log a_{H_2O}$$
(6)

- where *T* is temperature and *A* and *B* constants, with $A = \Delta H \div (2.303 \times R)$ and
- $B = \Delta S \div (2.303 \times R)$, given the ΔCp approximation.

Calibration of the thermometer

The new thermometer is calibrated from the relation between the logarithm of the equilibrium constant and 1/T, using the 161 low-P chlorite compositions (P < 4 kbar) and their estimated formation temperature according to the literature (Table 1). For those calculations, we assumed that $a\text{SiO}_2 = 1$, $a\text{H}_2\text{O} = 1$ and $\text{Fe}_{\text{total}} = \text{Fe}^{2+}$. From the regression analysis, we obtain the new linear equation:

289
$$\log K = -\frac{9400}{T(K)} + 23.40 \tag{7}$$

291 A quadratic regression yields the equation

293
$$\log K = \frac{11185729}{T(K)^2} - \frac{56598}{T(K)} + 72.3$$
 (8)

which provides a better fit in the range 300-350°C, but has no physical basis (cf. equation 6) and so should not be used outside the range 150-350°C. The relationship between reference temperatures (literature data) and temperatures calculated with the new thermometer for the calibration dataset is shown in Figure 2. The standard deviation from the 1:1 line is almost 46°C when using the combination of equations (7) and (8) (cf. Fig 2) and 65°C when using equation (7). However, the standard deviation calculation is biased by the 6 calculated temperatures exceeding 450°C (Fig 2) and is variable with T: in fact, the standard deviation is probably lower at low T (around ± 30 °C) and higher at T > 300°C (around ± 50 -60°C). This data scattering is similar to those obtained when testing the Inoue et al. (2009) and Vidal et al. (2006) models on the same literature dataset (XFe³+ estimated with the Vidal et al. 2006 model). Therefore, we consider that the correlation between estimated and expected T is satisfactory.

Testing the new geothermometer

Using equation (7) applied to the chlorite + quartz assemblage, the formation temperatures of Gulf Coast, Saint Martin and Toyoha chlorites were estimated assuming all iron is ferrous and 1 for H_2O activity. In fact, in all cases, the values of H_2O activity in the fluid are unknown and the assumption $aH_2O = 1$ seems to be reasonable in diagenetic to low-grade metamorphic conditions (Inoue et al. 2009), but remains locally questionable. The assumption $aSiO_2 = 1$ is justified in the case of quartz-bearing rocks.

The temperatures calculated with equation (7) are plotted (i) versus measured temperatures in Figure 3 for Gulf Coast chlorites, for which the high-spatial-resolution should ensures the best reliability of the analyses, (ii) versus Al^{IV} content in Figure 4 for Saint Martin and Toyoha chlorites assumed to be authigenic crystals. The temperatures obtained with the new thermometer range from 91 to 231°C, for an expected range of 102-232°C for the Gulf Coast samples. Few analyses (BHT~180°C) give too low a temperature (Figure 3) suggesting that they do not represent highest-*T* equilibrium composition. Generally, the measured and calculated temperatures are very close, with less than 20°C difference. For Saint Martin chlorites, the calculated temperatures range from 172 to 351°C (Figure 4) for an expected 220-340°C whereas for Toyoha chlorites, the temperatures range from 159 to 301°C for an expected 159-264°C (a single calculated temperature is outside the expected range). In all cases, the new equation appears to be highly reliable to obtain sensible temperature values, despite the assumption that all iron is ferrous and regardless of the rock-type considered.

Discussion

Suitability of chlorite analyses for calibration and testing

As pointed out by Hillier and Velde (1992), De Caritat et al. (1993), Jiang et al. (1994) and Essene and Peacor (1995), part of the compositional variations measured for low-T chlorites may be ascribed to contamination by kaolinite, or to interstratification with berthierine or smectite, which might also lead to erroneous structural formulae and an overestimation of vacancy contents (vacancies are then considered as an analytical artifact). The interstratification with smectite is assumed to be negligible here because only analyses with < 1 %wt total Na₂O + CaO + K₂O were selected, both for calibration and for testing, almost 90% of which actually have Na₂O + CaO + K₂O < 0.5% wt. The interstratification with

the 7 Å-phase berthierine, which is a common phenomenon in low-T chlorites (Xu and Veblen, 1996), is difficult to distinguish chemically from a true solid-solution without high-resolution structural control. In this respect, the use of a high-resolution analytical protocol (with AEM) for the Gulf Coast chlorites allows for avoiding the 7 or 10 Å-layer and quartz contaminations; it shows that octahedral vacancies in low-T chlorite are an actual feature and that their amount varies with T. Therefore, the deficit in octahedral cations (< 6 apfu) observed under all conditions (diagenetic or metamorphic) in the absence of significant smectite or mica contamination is real, and should be amplified in structural formula recalculation when accounting for Fe³⁺ content, which is not negligible for low-T chlorites (Beaufort et al. 1992).

An issue is the reliability of AEM data compared to EMP. Only three AEM databases were used here: Jahren and Aagaard (1992) and Lopez-Munguira et al. (2002) for calibration, and Gulf Coast chlorites analyses for testing. Lopez-Munguira et al. (2002) showed that the analyses obtained with both methods are highly similar. Bourdelle et al. (2012) also show the suitability of a FIB-AEM protocol to obtain chlorite analyses with a level of confidence similar to EMP analyses: this is the protocol used here to analyse the Gulf Coast chlorites. On this basis, we consider that the reliability of the analyses used for calibration or testing is not affected by the applied analytical method.

In a broader perspective and for practical purposes, we aimed at the largest possible database for calibration, offering a wide coverage of geological parameters, like lithology, temperature gradient, kinetics, fluid regimes, oxidation state, chlorite composition (including XFe³⁺), chlorite precursors, impurities and analytical contaminations (insofar as they are of limited extent). This was considered the best way to limit the effect of each parameter, with the aim of a general thermometer for low-T chlorites. Therefore, all the data used for thermometer calibration were considered as suitable, provided they met the imposed chemical criteria. As a result, the database used in this study is the largest ever gathered for low-T chlorite thermometry, with 161 analyses for $T < 350^{\circ}$ C, P < 4kbar, of which 94 analyses are for $T < 250^{\circ}$ C, compared to 23 analyses (for $T < 350^{\circ}$ C) for Inoue et al. (2009; from Vidal et al. 2001 appendix), with 11 analyses for $T < 250^{\circ}$ C.

Comparison with recent chlorite thermometers

The results obtained on Gulf Coast, Saint Martin and Toyoha chlorites with the new thermometer (using the linear equation (7) and the assumption $Fe_{total} = Fe^{2+}$) are compared in Figure 5 to the two main non-empirical models (Vidal et al. 2005, 2006; Inoue et al. 2009) for

which Fe³⁺ content is taken into account. Note that many analyses were automatically excluded in Vidal's model by their Si content higher than 3 apfu. The thermodynamic model of Walshe (1986) was not considered, as it does not allow for Si contents lower than 3 apfu.

For the Gulf Coast chlorites, Vidal et al. (2005, 2006) and Inoue et al. (2009) models give substantially similar results, with a range of formation temperatures of 84-317°C and 113-325°C, respectively (Figure 5a). The two sets of results are very similar (Figure 5a, inset) and are very close to those obtained with equation (7). However, some analyses near ~200°C branch off the 1:1 trend, yielding overestimated temperatures with Inoue and Vidal models. These analyses are characterised by high Al content and low XFe³⁺ (according to Vidal procedure) in contrast to the high-*T* analyses of the 1:1 trend, which have low Al content and high XFe³⁺ (~35-40%). Important is that, regardless of these compositional variations including XFe³⁺, the present thermometer, unlike the two others, yields a single trend of temperatures for these two groups, consistent with measured temperatures (cf. Figure 3).

For the Saint Martin chlorites, Vidal et al. (2005, 2006) and Inoue et al. (2009) models yield results in the ranges 50-321°C and 172-313°C, respectively. In this case, most of the temperatures obtained with Vidal's model are underestimated and differ from the two other thermometers, whereas the newly proposed equation (7) gives temperatures that are very similar to those obtained with Inoue's model.

For Toyoha chlorites, the situation is similar to the Saint Martin case, with a scatter of results obtained with Vidal's thermometer (70-265°C), and results of the two other thermometers close to each other and in better agreement with the expected range (159-264°C).

For each of the three geological fields, the results are similar and in the expected temperature windows for Inoue's and the newly proposed model, whereas Vidal's model gives more disparate, often underevaluating results. These differences in calculated temperatures can be explained by the choices of end-members (and site assignment of iron) and of calibration basis (more weight of the high-T, low-Fe³⁺ analyses in Vidal and Inoue databases). The excellent agreement between Inoue and Vidal's models in Figure 5a (inset) in contrast to Figure 5b and c (insets) is not fortuitous as this is the sole case for which XFe³⁺ input ratios are obtained with Vidal et al. (2006) iterative method. In summary, the comparison of the new thermometer and Vidal and Inoue's models shows that equation (7), based on an ordered distribution and the Fe_{total} = Fe²⁺ assumption, performs well in the case of low-T chlorites.

Choice of a semi-empirical ordered model for low-T chlorites

The semi-empirical approach, based on a reaction, appears as a good alternative solution between a purely thermodynamic treatment, which requires knowledge of the many end-member and mixing properties (as Vidal et al. 2006), and the purely empirical approach, which does not consider the bulk composition (as Cathelineau 1988). As regards the cationic distribution, Inoue et al. (2009) suggest that one reason for the inaccuracy in low-*T* domain of the Vidal et al. (2001) model, rearranged by Vidal et al. (2005, 2006), is the ordered site occupancy. This statement is derived from the polytype distribution (Ia, Ib, IIb) according to the metamorphic grade. However, in the absence of univocal information, in particular on the validity of each cationic distribution model according to polytype, we prefer to base the new thermometer on a semi-ordered distribution. The results presented above show that it gives at least as reliable estimates as Inoue's disordered model for the three geological fields of Gulf Coast, Saint Martin and Toyoha. The cation distribution model is therefore not the main issue for chlorite thermometry. On the basis of the above comparisons, it appears that the calibration (*T* range and compositional space) and the choice of end-members determine the pertinence of the various models.

Testing the effect of non-ideality

As done by Inoue et al. (2009), the non-ideal contribution of the site mixing was neglected in the present study. This assumption was required to calibrate a semi-empirical T = f(K) geothermometer, but it needs to be evaluated, considering that sudoite and trioctahedral chlorite may coexist in equilibrium in a few cases (in low-T, low-P or low-T, high-P aluminous metapelites, e.g. Theye et al. 1992). For the assumption of ideal mixing to be tenable, one must verify that (i) for a given chlorite composition, the contribution of the non-ideality varies linearly with 1/T, and (ii) at given T, the variation of the non-ideal contribution with chlorite composition is small, so that the net effect of non-ideality is grossly linear in T and so makes a linear calibration possible.

For the first point, the expression of the non-ideality is:

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$$K = \prod_{j} (a_{ideal} \cdot \gamma)_{j}^{v_{j}} \text{ with } \gamma_{j} = \prod_{s} \prod_{m} \gamma_{m}^{n_{s}}$$
 (9)

where, for a j component, γ is the non-ideality coefficient, n_s is the multiplicity of the site s and m are the relevant cations. The log/ln K can be expressed as the sum of an ideal and a non-ideal part:

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$$R.T. \ln K = R.T. \ln \left(\prod_{j} a_{ideal} \right)_{j} + R.T. \ln \left(\prod_{j} \gamma \right)_{j}$$
 (10)

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Neglecting ternary and quaternary interactions, the non-ideal contribution may be written:

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$$n_s.R.T. \ln \gamma_m = \sum W_{ij}.X_i.X_j. \left[\frac{Q_m}{X_m} - 1 \right]$$
 (11)

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where W_{ij} are the Margules parameters, X_m is the mole fraction of cation m and Q_m is the number of i, j subscripts that are equal to m. Clearly, from this formalism, the non-ideal contribution in the log/ln K calculation is linear in 1/T for a fixed composition.

For the second point, Bourdelle (2011) tested the new model for various chlorite compositions and P-T conditions. Starting with an average composition and an approximation of Margules parameters defined by Vidal et al. (2001, 2005, 2006), he used equations (10) and (11) and the defined semi-ordered distribution to calculate the non-ideal contribution for several temperatures at given P, considering symmetric binary interactions. The author introduced variation in the composition and conditions, by varying the XSi_{T2}, XMg (ratio), $X\square$ and pressure. The results of these tests are plotted in Figure 6 and show, in the range of 100-350°C and neglecting the W(M2+M3) parameters (following Vidal et al. 2005, 2006) as well as the pressure factor, that the variations of XMg ratio affect only very slightly the variation of $\log/\ln K$ with 1/T. This holds especially if the tested chlorite compositions are only considered in the range of temperatures to which they refer (i.e. the Si-rich composition, Si = 3.4 apfu, mainly characterises very low-T chlorites, contrary to the Si-poor composition, Si = 2.6 apfu). The non-ideal contribution mainly affects the y-intercept of the In K = f(1/T)curve, which does not preclude a linear calibration. Thus, we conclude that the variation of the non-ideal contribution with composition (in the classical range of diagenetic chlorites) is sufficiently small for the linearity of $\log/\ln K$ in 1/T to be grossly preserved. Therefore, for the P-T domain and the composition range investigated here, calibration on a linear basis is justified and, actually, any effect of non-ideality is implicitly taken into account in the thermometer equation.

Link between Fe^{3+} content, octahedral vacancies, thermometer calibration and temperature calculation

The main compositional features of chlorite evolution with increasing T are long recognized as the Tschermak substitution with a contribution of di-trioctahedral substitution at low T. This decrease of octahedral vacancies with increasing T is therefore built in any calibration database, particularly when the database includes lower-T chlorites. Besides, accounting for Fe³⁺ in the structural formula recalculation on a fixed oxygen basis arithmetically reduces the number of cations, and implies an increase of octahedral vacancies, which in turn links indirectly Fe³⁺ content and temperature in any thermometric formulation. However, this effect of Fe³⁺ remains subordinate with respect to the main compositional evolutions with T mentioned above (compare Figures 6a and 6b in Inoue et al. 2009). Accordingly, variations of the chlorite + quartz $\log K$ remain negatively correlated with 1/T, regardless of the assumption made on the oxidation state of iron. This justifies calibrating a chlorite thermometer on one or the other assumption, i.e. $Fe_{total} = Fe^{2+}$ or real Fe^{3+} values. A calibration based on the latter choice should lead to higher precision, but calibration based on the former case may be a good trade-off of practicability and simplicity (Fe³⁺ content not required) against precision. Admittedly, the sudoite activity is poorly evaluated, i.e. biased by the non consideration of Fe³⁺ content, but this possible bias is built in the calibration. The best demonstration of the validity of this trade-off is the success of the thermometer presented here in its applications to various geological contexts.

Conclusion

Various concerns have been raised on the relevance, applicability and precision of chlorite thermometry (e.g. De Caritat et al. 1993; Jiang et al. 1994; Essene and Peacor 1995). However, recent developments (e.g. Vidal et al. 2001, 2005, 2006; Inoue et al. 2009) have led to a reappraisal of this technique and to new opportunities in exploiting the information contained in chlorite composition. Comparative studies (e.g. Inoue et al. 2009; this paper) have yet pointed out that order-disorder, compositional space, non-ideality and Fe^{3+} effects may be limitations and/or sources of disagreement. The approach developed here, compensating these sources of variability through the use of a large database for linear calibration, proves successful in its application to low-T chlorites. This success may lie in the fact that the linear dependence of log K with 1/T either integrates these effects (were they

non-ideality or Fe³⁺ content), or is not significantly affected by them, the backbone of the variation being the combined effects of the Tschermak and di-trioctahedral substitutions.

The new geothermometer allows one to account for all the low-T chlorite compositions, especially Si-rich compositions that characterize diagenetic chlorites. The comparison of the new thermometer with the recent ones that require Fe³⁺ content estimation shows that it is possible to obtain reliable results (\pm 20°C in most cases) without any measurement or assumption on the Fe³⁺ content. In this respect, the present semi-empirical thermometer is a much practical tool, well suited for, e.g., the handling of large analytical datasets in exploration geology. However, it should be kept in mind that the new equation proposed here is only valid over its calibration range ($T < 350^{\circ}$ C, P < 4 kbar) and that any use out of these limits, in particular in the metamorphic realm, is not recommended – mainly because of the assumptions on the heat-capacity and volume changes of the reaction, the pressure effect and the water activity in the new model, which may no longer be valid. Above 350°C, the thermometry of Vidal et al. (2005, 2006) remains probably the most convenient approach.

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Tables

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Table 1 Published data and methods to estimate temperature for the analyses that have been used to calibrate the new thermometer

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Study	Temperature estimation methods					
Boles and Franks (1979)	in situ temperature measurement					
McDowell and Elders (1980)	in situ temperature measurement					
Cathelineau (1988)	in situ temperature measurement, data compilation					
Hutcheon (1990)	in situ temperature measurement					
Bevins et al. (1991)	chl-thermometry, mineral assemblages					
Jahren and Aagaard (1992)	in situ temperature measurement					
De Caritat et al. (1993)	data compilation, chl-thermometry, carbonates thermometry					
Rahn et al. (1994)	chl-thermometry, vitrinite reflectance, fluid inclusions					
Schmidt et al. (1997)	chl-thermometry, fluid inclusions, isotopy ($\Delta^{18}O_{\text{qz-calcite}}),$ IC and CC index					
Xie et al. (1997)	chl-polytypism, chl-thermometry					
Lopez-Munguira et al. (2002)	chl-thermometry					
Mas et al. (2006)	in situ temperature measurement					
Koroknai et al. (2008)	chl-thermometry, mineral assemblages					

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Table 2 AEM analyses of Gulf Coast chlorites (crystal rims analyses). Elements contents are given in atom per formula unit (O = 14 apfu). All iron is considered as ferrous

Campla	AZ#159	AZ#159	AZ#159	AZ#159	ST#470	ST#470	ST#470	CK#2	CK#2	CK#2
Sample	9230	9230	9230	9230	10717	10717	10717	11924	11924	11924
Analysis	chl10	chl17	chl30	chl28	chl27	chl29	chl30	chl47	chl50	chl54
BHT (°C)	102	102	102	102	121	121	121	129	129	129
BHP (bars)	300	300	300	300	590	590	590	660	660	660

	Si	2.98	2.95	2.95	2.91	3.08	3.19	3.01	2.96	2.90	2.91
	Ti	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.00
	Al	2.76	2.76	2.75	2.86	2.55	2.60	2.77	2.90	2.98	2.86
	Fe^{2+}	2.18	2.20	2.34	2.46	2.81	2.55	2.53	2.45	2.37	2.53
	Mn	0.00	0.01	0.02	0.00	0.02	0.02	0.01	0.00	0.01	0.00
	Mg	1.65	1.71	1.60	1.39	1.15	1.11	1.22	1.18	1.31	1.29
	Ca	0.03	0.01	0.00	0.00	0.01	0.00	0.02	0.01	0.00	0.01
	Na	0.00	0.00	0.01	0.03	0.00	0.03	0.00	0.02	0.00	0.00
	K	0.05	0.05	0.06	0.05	0.07	0.05	0.06	0.06	0.06	0.07
667											
		CK#2	CK#2	CK#2	CK#2	LA#1	LA#1	LA#1	LA#1	ST#356	ST#356
	Sample	12196	12196	12196	12196	13559	13559	13559	13559	14501	14501
	Analysis	chl20	chl18	chl19	chl21	chl23	ch120	chl11	chl17	chl36	ch134
	BHT (°C)	135	135	135	135	149	149	149	149	166	166
	BHP (bars)	690	690	690	690	850	850	850	850	800	800
	Si	2.96	3.05	3.02	3.02	2.88	2.86	2.83	2.78	2.84	2.71
	Ti	0.00	0.02	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00
	Al	2.85	2.83	2.91	2.81	2.91	2.92	2.95	3.07	2.85	3.00
	Fe ²⁺	2.45	2.38	2.38	2.39	3.28	3.36	3.41	3.28	2.05	2.07
	Mn	0.00	0.02	0.00	0.01	0.01	0.00	0.00	0.03	0.00	0.00
	Mg	1.34	1.18	1.20	1.30	0.53	0.48	0.47	0.49	1.96	1.98
	Ca	0.00	0.04	0.01	0.01	0.02	0.00	0.01	0.03	0.00	0.00
	Na	0.00	0.00	0.00	0.00	0.02	0.00	0.00	0.03	0.03	0.00
	K	0.00	0.03	0.01	0.08	0.03	0.04	0.04	0.02	0.03	0.05
668		0.00	0.02	0.01	0.00	0.03	0.01	0.01	0.03	0.03	
000											
	Sample	CW#1	CW#1	CW#1	CW#1	WR#C1	WR#C1	WR#C1	FR#1	FR#1	FR#1
		14277	14277	14277	14277	17805	17805	17805	18946	18946	18946
	Analysis	chl13	chl14	chl32	chl31	chl13	chl18	chl16	chl26	chl30	chl28
	BHT (°C)	191	191	191	191	191	191	191	204	204	204
	BHP (bars)	750	750	750	750	1050	1050	1050	1150	1150	1150
	Si	2.68	2.68	2.65	2.63	2.98	2.97	2.90	2.88	2.87	2.87
	Ti	0.00	0.02	0.01	0.01	0.00	0.00	0.00	0.00	0.00	0.01
	Al	3.38	3.22	3.24	3.21	3.00	2.91	2.95	2.64	2.62	2.60
	Fe^{2+}	2.68	2.98	3.00	3.01	2.38	2.39	2.48	2.01	2.08	2.07
	Mn	0.20	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.00
	Mg	0.82	0.79	0.80	0.85	1.08	1.26	1.28	2.24	2.19	2.23
	Ca	0.05	0.00	0.00	0.04	0.01	0.02	0.00	0.00	0.02	0.01
	Na	0.00	0.00	0.07	0.00	0.03	0.00	0.01	0.00	0.00	0.00
	K	0.04	0.04	0.00	0.03	0.03	0.05	0.02	0.05	0.06	0.07
669											
	C 1	FR#1	AL#1	_							
	Sample	18946	19110	19110	19110	19110	20711	20711	20711		
	Analysis	chl29	chl17	chl18	chl27	chl25	chl31	chl28	chl29	_	
	-										

BHT (°C)	204	216	216	216	216	232	232	232
BHP (bars)	1150	1150	1150	1150	1150	1200	1200	1200
Si	2.84	2.66	2.64	2.66	2.61	2.94	2.88	2.85
Ti	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.00
Al	2.64	3.21	3.21	3.13	3.16	2.52	2.60	2.64
Fe^{2+}	2.15	2.47	2.53	2.43	2.44	2.74	2.95	2.93
Mn	0.00	0.02	0.00	0.00	0.00	0.00	0.00	0.00
Mg	2.11	1.38	1.32	1.49	1.55	1.50	1.36	1.36
Ca	0.01	0.01	0.02	0.01	0.00	0.00	0.00	0.00
Na	0.02	0.00	0.00	0.03	0.00	0.02	0.00	0.02
K	0.07	0.01	0.07	0.04	0.05	0.06	0.07	0.06

Table 3 Ideal activities of solid-solution end-members used in the calculation of the three chlorite geothermometers

End-members (abbreviations)	Chemical formula* (half formula unit)	Equilibrium / Ideal activities [†]				
Vidal et al. (2005, 2006) mo	del					
Thermodynamic formalism		2 Clin + 3 Mg-Sud = 4 Mg-Am + 7 Qtz + 4 H2O				
Clinochlore (Clin)	$(AlMg_5)(Si_3Al)$	$=4\;(X_{Mg,M1})(X_{Mg,M2+M3})^4(X_{Si,T2})(X_{Al,T2})$				
Daphnite (Daph)	$(AlFe_5)(Si_3Al)$	$=4\ (X_{Fe,M1})(X_{Fe,M2+M3})^4(X_{Si,T2})(X_{Al,T2})$				
Mg-Amesite (Mg-Am)	$(Al_2Mg_4)(Si_2Al_2)$	$= 64 \; (X_{\square,M1})(X_{Al,M2+M3})^2 (X_{Mg,M2+M3})^2 (X_{Si,T2})(X_{Al,T2})$				
Fe-Amesite (Fe-Am)	$(Al_2Fe_4)(Si_2Al_2)$	$= 4 \; (X_{\square,M1})(X_{Al,M2+M3})^2 (X_{Mg,M2+M3})^2 (X_{Si,T2})(X_{Al,T2})$				
Mg-Sudoite (Mg-Sud)	$(Al_3Mg_2)(Si_3Al)$	$= (X_{Al,M1})(X_{Mg,M2+M3})^4(X_{Al,T2})^2$				
Inoue et al. (2009) model						
$T(K) = 1/(0.00293 - 5.13 \times 10^{-4} \times 10^{-4})$	$\log K + 3.904 \times 10^{-5} \times (\log K)^2$	$Mg\text{-Chl }S+3\ Mg\text{-Sud}=3\ Mg\text{-Am}+7\ Qz+4\ H_2O$				
Mg-Chlorite Sa (Mg-Chl S)	$(Mg_6)(Si_4)$	$= (X_{Mg,oct})^6 (X_{Si,tet})^2$				
Daphnite ^b (Daph)	$(AlFe_5)(Si_3Al)$	= 59.720 $(X_{Fe,oct})^5(X_{Al,oct})(X_{Si,tet})(X_{Al,tet})$				
Mg-Sudoite (Mg-Sud)	$(Al_3Mg_2)(Si_3Al)$	$=1728\;(X_{Mg,oct})^2(X_{Al,oct})^3(X_{\square,oct})(X_{Si,tet})(X_{Al,tet})$				
Mg-Amesite ^c (Mg-Am)	$(Al_2Mg_4)(Si_2Al_2)$	$=45.563~(X_{Mg,oct})^4(X_{Al,oct})^2(X_{Al,tet})^2$				
This study model						
$T(K) = 9400/(23.40 - \log K)$		Mg -Chl S + 3 Mg -Sud = 3 Mg -Am + 7 Qz + 4 H_2O				
Mg-Chlorite S (Mg-Chl S)	$(Mg_6)(Si_4)$	$= (X_{Si,T2})^2 (X_{Mg,M2+M3})^4 (X_{Mg,M1+M4})^2$				
Fe-Chlorite S (Fe-Chl S)	$(Fe_6)(Si_4)$	$= (X_{Si,T2})^2 (X_{Fe,M2+M3})^4 (X_{Fe,M1+M4})^2$				
Mg-Amesite (Mg-Am)	$(Al_2Mg_4)(Si_2Al_2)$	$= (X_{AI,T2})^2 (X_{Mg,M2+M3})^4 (X_{AI,M1+M4})^2$				
Fe-Amesite (Fe-Am)	$(Al_2Fe_4)(Si_2Al_2)$	= $(X_{Al,T2})^2 (X_{Fe,M2+M3})^4 (X_{Al,M1+M4})^2$				
Mg-Sudoite (Mg-Sud)	$(Al_3Mg_2)(Si_3Al)$	$= 256 \ (X_{Si,T2})(X_{Al,T2})(X_{Al,M1+M4}) \\ (X_{\square,M1+M4})(X_{Mg,M2+M3})^2(X_{Al,M2+M3})^2$				

Fe-Sudoite (Fe-Sud)	$(Al_3Fe_2)(Si_3Al)$	$= 256 (X_{Si,T2})(X_{Al,T2})(X_{Al,M1+M4})$
		$(X_{\Box,M1+M4})(X_{Fe,M2+M3})^2 (X_{Al,M2+M3})^2$

Note: $X_{i,s}$ is the mole fraction of the *j* cation on the *s* site

Equivalents in Inoue et al. (2009): ^a Al-free chlorite, ^b chamosite, ^c corundophilite. * Structural formula based on

O₁₀(OH)₈. † Ideal activities are completed with a non-ideal contribution in the Vidal et al. (2005, 2006) model

Table 4 Chlorite solid-solution end-members and cationic site repartition used in the new geothermometer calculation

End-members	T1(2)	T2(2)	†	(M2+M3)(4)	†	(M1+M4)
Mg-Chlorite S	Si(2)	Si(2)		Mg(4)		Mg(2)
Fe-Chlorite S	Si(2)	Si(2)		Fe(4)		Fe(2)
Mg-Sudoite	Si(2)	SiAl		Al(2)Mg(2)		Al□
Fe- Sudoite	Si(2)	SiAl		Al(2)Fe(2)		Al□
Mg-Amesite	Si(2)	Al(2)		Mg(4)		Al(2)
Fe-Amesite	Si(2)	Al(2)		Fe(4)		Al(2)
Mg			4	$Mg-Mg_{M1} \\$	3	$(Fe_{M1}+Mg_{M1})\;x\;XMg$
Fe			4	$Fe-Fe_{M1}$	3	$\left(Fe_{M1}+Mg_{M1}\right)x\;XFe$
Fe + Mg			2	$4 - (Al^{VI} - Al^{IV})$	2	2 - $(Al_{M1+M4} + \Box)$
Al		4-Si	1	Al ^{VI} - Al ^{IV}	1	$\mathrm{Al^{IV}}$
				-	1	$(Al^{VI} - Al^{IV})/2$

[†] indicate the sequence in which the cation assignment is made

Figures

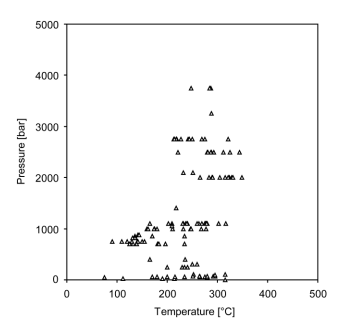


Fig. 1 *P-T* data for 161 chlorites analyses compiled from literature (Table 1)



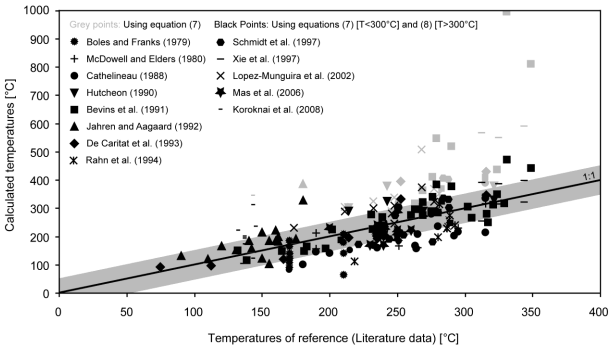


Fig. 2 Relationship between temperatures of reference (literature data) and temperatures calculated with the new thermometer (161 analyses used for thermometer calibration, listed in Table 1), considering the chemical composition of chlorites formed at P < 4 kbar. Black symbols: temperatures calculated with the linear equation (7) and the quadratic equation (8) when T > 300°C. Grey symbols: temperatures calculated with the linear equation (7) only.

The shaded zone indicates the 1:1 line $\pm 50^{\circ}$ C. See text for details

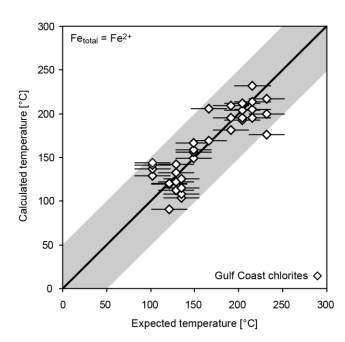
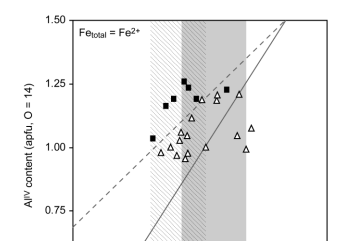


Fig. 3 Comparison of measured/expected temperatures (BHT \pm 20°C) for chlorites of the Gulf Coast formation versus temperatures calculated using the new geothermometer (all Fe is ferrous)

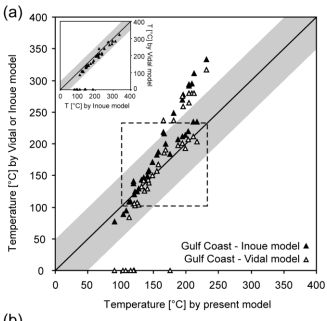


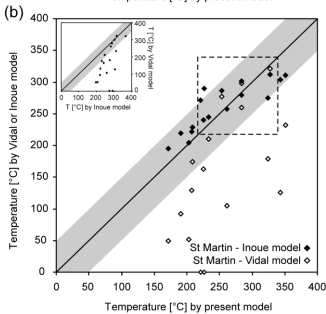
Temperature by present model [°C]

0.50 +

Fig. 4 Plot of tetrahedral Al contents versus calculated temperatures using the new geothermometer (all Fe is ferrous) for Saint Martin and Toyoha chlorites. The range of expected temperatures is indicated by the shaded area for St Martin chlorites, by the hatched area for Toyoha chlorites. The solid and dashed curves are the equations of Cathelineau (1988) and Hillier and Velde (1991) empirical thermometers, respectively

Saint Martin △ Toyoha ■





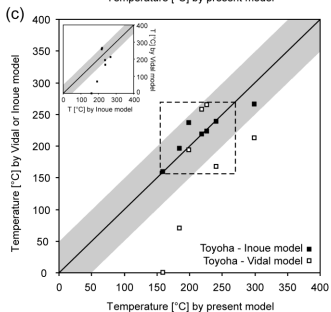


Fig. 5 Comparison of calculated temperatures using Inoue et al. (2009), Vidal et al. (2006) and the present chlorite geothermometers. Solid and open symbols refer to temperatures calculated by Inoue et al. (2009) and Vidal et al. (2006) models, respectively. Dashed rectangular areas correspond to the expected temperature ranges for the Gulf Coast (a), Saint Martin (b) and Toyoha (c) datasets. The shaded zone indicates the 1:1 line $\pm 50^{\circ}$ C.

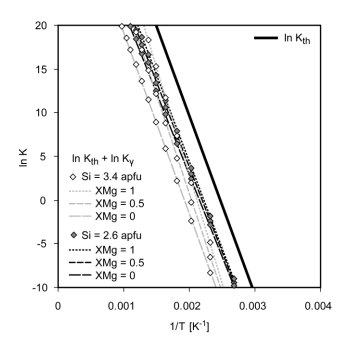


Fig. 6 Effect of non-ideality on $\ln K$, considering symmetric binary interactions, $X \square = 0.2$ apfu, W(M2-M3) = 0 and various Si content and XMg ratio; modified after Bourdelle (2011). The solid curve corresponds to the thermometer calibration (noted $\ln K_{th}$), which is indicated as reference and is calculated according to 1/T (not from the fixed compositions). The variations of XMg ratio affect only very slightly the slope of the $\ln K = f(1/T)$ curve, especially for the Si-poor composition case