

Fluid circulation along an oceanic detachment fault: insights from fluid inclusions in silicified brecciated fault rocks (Mid-Atlantic Ridge at 13°20'N)

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1	Fluid circulation along an oceanic detachment fault: insights from fluid
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4	A. Verlaguet ^{1*} , D. Bonnemains ² , C. Mével ² , J. Escartín ^{2,3} , M. Andreani ⁴ , F.
5	Bourdelle ⁵ , M-C. Boiron ⁶ , V. Chavagnac ⁷
6	, , , , , , , , , , , , , , , , , , , ,
7 8	¹ Sorbonne Université, CNRS-INSU, Institut des Sciences de la Terre, ISTeP UMR 7193, F 75005 Paris, France
	² Marine Geosciences Group, IPGP CNRS UMR 7154, Paris, France
9	•
10	³ Laboratoire de Géologie, UMR 8538, Paris, France
11	⁴ Laboratoire de Géologie, Université de Lyon, Lyon, France
12	⁵ Univ. Lille, IMT Lille Douai, Univ. Artois, Yncrea Hauts-de-France, ULR 4515 - LGCgE,
13	Laboratoire de Génie Civil et géo-Environnement, F-59000 Lille, France
14	⁶ Université de Lorraine, CNRS, GeoRessources, F-54000 Nancy, France
15	⁷ Géosciences Environnement Toulouse (GET), Université Paul Sabatier Toulouse 3, CNRS UMR 5563,
16	IRD, Toulouse, France
17	
18 19	* Corresponding author: anne.verlaguet@sorbonne-universite.fr
13	Corresponding author: aime.verraguetta/sorbonne-universite.ir
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22	Keypoints
23	• MAR 13°20'N corrugated detachment fault is composed of pervasively silicified mafic
24	breccias overplated from hangingwall diabases
25	• Quartz fluid inclusions record mixing of hangingwall silica-rich brines with footwall
26	serpentinite-derived fluids (H ₂ ±CH ₄) in detachment
	/
27	• This heterogeneous fluid circulation in shallow detachment fault is inconsistent with
28	models of detachments channeling deep fluid flow
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Abstract

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The MAR 13°20'N corrugated detachment fault is composed of pervasively silicified mafic cataclastic breccias, instead of ultramafics and gabbros commonly found at other detachments. These breccias record overplating of hangingwall diabases, with syntectonic silicification due to important influx of silica-iron-rich fluids, able to leach alkalis and calcium. Fluids trapped in quartz inclusions show important salinity variations (2.1-10 wt.% NaCl eq.) indicating supercritical phase separation. Fluid inclusions also contain minor amounts of H₂+CO₂+CH₄+H₂S, with high H₂/CO₂ and H₂/H₂S ratios, signatures typical of ultramafic-hosted vent fluids. We propose that seawater infiltrated the hangingwall upper crust at the axis adjacent to the active detachment, reaching a reaction zone at the dyke complex base (~2 km). At >500°C, fluids become Si-rich during diabase alteration (amphibolite-facies alteration in clasts), and undergo phase-separation. Brines, preferentially released in the nearby detachment fault during diabase brecciation, mix with serpentinite-derived fluids bearing H₂ and CH₄. Cooling during detachment deformation and fluid upward migration triggers silica precipitation at greenschist-facies conditions (quartz+Fe-richchlorite+pyrite). Important variations in fluid inclusion salinity and gas composition at both sample and grain scales record heterogeneous fluid circulation at small spatial and short temporal scales. This heterogeneous fluid circulation operating at <2 km depth, extending both along-axis and over time, is inconsistent with models of fluids channeled along detachments from heat sources at the base of the crust at the fault root. Present-day venting at detachment footwall, including Irinovskoe, is instead likely underlain by fluid circulation within the footwall, with outflow crossing the inactive detachment fault near-surface.

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Keywords

- 53 MAR 13°20'N, silicified detachment, fluid inclusions, mafic breccias, phase separation,
- 54 hangingwall overplating

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Plain language summary

Here we present constraints on fluid circulation along the 13°20′N oceanic detachment fault along the Mid-Atlantic Ridge. Rocks recovered in situ with a deep-sea robot yield mafic breccias, instead of serpentinized mantle rocks commonly found at other detachments. They likely originate from the base of the hangingwall dyke complex, brecciated during fault exhumation. These rocks are intensely silicified (quartz mineralization), resulting from upflow circulation of silica-rich fluids derived from reactions with mafic rocks in a reaction zone. Fluid inclusion (micrometric cavities in quartz crystals that trapped circulating fluid) analyses reveal highly-saline fluids likely formed by phase separation, while traces of hydrogen and methane likely record serpentinization. We thus propose that seawater infiltrated the crust down to a reaction zone at its base (2 km depth), where it became silica-rich by rock hydrothermal alteration. Upon brecciation, these silica-rich brines were released in the detachment where they mixed with fluids coming from footwall rock alteration. Temperature and pressure drops during fluid upflow promoted intense quartz crystallization. The active Irinovskoe hydrothermal site, sitting on the detachment fault ~5 km off-axis, is unrelated to fluid circulation in the detachment plane, and likely linked to a heat source within the footwall and directly below it.

1. Introduction

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Low-angle detachment faults are common along slow- and ultra-slow spreading ridges, forming primarily at ridge sections with reduced melt supply (Buck et al., 2005; Cannat et al., 2006; Escartín et al., 2008; Tucholke et al., 2008). Rooting deep below the rift valley floor, detachments can operate over long periods of time (up to a few Myrs), exhume deep-seated materials from the oceanic lithosphere, and lead to the formation of oceanic core complexes (OCCs). OCCs are often capped by a fault surface displaying corrugations parallel to spreading. Ridge sections hosting OCCs tend to be associated with high seismicity rates relative to magma-rich, symmetrical ridge sections, and often display active hydrothermal venting (Escartín et al., 2008; Son et al., 2014). The microseismicity along the Northern Atlantic and the Southwest Indian Ridge detachments (deMartin et al., 2007; Parnell-Turner et al., 2017; Tao et al., 2020) reaches sub-Moho depths (between ~8 and ~13 km bsf). These results suggest that detachments are associated to a thick lithosphere, and that brittle deformation may provide potential pathways for fluid circulation reaching deep levels. Indeed, fault zone rocks from various OCCs along the MAR display evidence for fluid-rock interactions coeval with deformation. In most cases, the fault zone is characterized by deformed ultramafics and gabbros recrystallized into talcschists and amphibolites, respectively (Boschi et al., 2006; Escartín et al., 2003, 2017; Karson et al., 2006; MacLeod et al., 2002; McCaig et al., 2007; Schroeder & John, 2004). Moreover, active hydrothermal fields are often found on the OCC surface, with hydrothermal circulation crosscutting pre-existing and/or inactive fault zones, as observed at Rainbow (Andreani et al., 2014), Ashadze (Ondréas et al., 2012), Logatchev (Petersen et al., 2009), Lost City (Fruh-Green et al., 2003) or Von Damm (Hogkinson et al., 2015). To date, however, there is still limited information regarding fluid sources and pathways along the detachment fault zone at depth, the location and nature of heat sources animating this hydrothermal circulation, and the possible links between the hydrothermal activity observed at the surface of these OCCs and the flow along the detachment.

The 13°20'N detachment fault, located on the western flank of the Mid-Atlantic Ridge (MAR), exposes a structurally continuous and corrugated detachment fault surface (e.g, Escartín et al., 2017; MacLeod et al., 2009; Smith et al., 2006). While morphology is similar to OCCs elsewhere (Parnell-Turner et al., 2018), sampling of the detachment fault zone reveals unique characteristics. Indeed, while other detachment faults are composed of deformed and hydrated footwall ultramafics and gabbros (Boschi et al. 2006; Escartín et al., 2003, 2016; Karson et al., 2006; MacLeod et al.,

2002; McCaig et al., 2007; Schroeder & John, 2004), the well-preserved and corrugated 13°20'N detachment fault zone is composed essentially of highly silicified mafic cataclastic breccias (Bonnemains et al., 2017; Escartín et al., 2017). These silicified fault breccias likely result from the overplating of mafic material from the base of the hangingwall dyke complex into the footwall during fault exhumation (Bonnemains et al., 2017); these rock types have not been reported from any other studied OCC. The silicified fault zone is ~70 m or thicker, and the mechanisms of strain localization and fault development leading to a corrugated structure seem to operate independently of lithologies within the fault zone (e.g., Parnell-Turner et al., 2018). Fault zone silicification is evidenced by massive quartz precipitation that may result from elevated fluxes of silica-rich fluids, possibly syntectonic (Bonnemains et al., 2017). This detachment fault surface also hosts the active Irinovskoe hydrothermal site, where several black smokers discharge high temperature fluids (Escartín et al., 2017; MacLeod et al., 2009).

The 13°20'N silicified fault rocks, and the fluid inclusions trapped in quartz crystals, provide a unique opportunity to investigate the nature, sources and pathways of fluids circulating within an active detachment fault zone, and to compare them to fluid circulation feeding the active footwall Irinovskoe hydrothermal site. Indeed, to date geochemical and fluid inclusion studies have been conducted on hydrothermally altered rocks from both hangingwall (e.g., Delaney et al., 1987; Humphris et al., 1998; McCaig et al., 2007, 2010; Tivey et al., 1998; Vanko et al., 2004) and footwall of detachments (e.g., Andreani et al., 2014; Boschi et al., 2006; Castelain et al., 2014). However, in these either the link to detachment fault deformation is not established, or the associated hydrothermal systems post-date detachment activity, as in active and inactive systems preserved at the footwall of oceanic detachments (e.g., Andreani et al., 2014; Escartín et al., 2017; Hodgkinson et al., 2015; Ondréas et al., 2012). Thus, studies addressing syntectonic fluid flow at detachments are scarce, and rely on geochemical data from rock samples, (e.g., McCaig et al., 2007, 2010), not from fluids.

To constrain fluid-rock interactions within an active detachment fault zone, here we present a study of fluid inclusions trapped in quartz crystals from silicified fault rocks of the 13°20'N OCC. A microthermometric study was conducted on >100 fluid inclusions from four representative fault rock samples collected *in situ* at different outcrops throughout the 13°20'N fault zone, and with varying degrees of silicification. Fluid inclusion composition was also investigated by Raman spectroscopy. Whole-rock geochemical analyses were performed to constrain nature and

composition of the host rocks, and chlorite analyses to constrain the temperature of quartz-chlorite crystallization. With these results and available geological constraints, we propose a model of hydrothermal fluid circulation within an active oceanic detachment fault.

2. Geological background of 13°20'N OCC and sampling

2.1. Geological setting

The 13°20'N oceanic detachment displays a structurally continuous fault surface with prominent extension-parallel corrugations, exposed at the seafloor, and that roots at the rift valley floor. This OCC develops on the western flank of the MAR, which spreads at a full rate of 24.6 mm/yr (MacLeod et al., 2009; Smith et al., 2008), and is likely active based on its morphology, the absence of late tectonic disruption (Escartín et al., 2017; MacLeod et al., 2009; Mallows & Searle, 2012), and seismic activity. Microseismicity defines a curved fault plane reaching >10 km below the ridge axis (Parnell-Turner et al., 2017), reminiscent of that of other detachments such as TAG (deMartin et al., 2007) or Longqi (Tao et al., 2020). This OCC and its detachment fault were extensively investigated and sampled during the ODEMAR cruise, using deep-sea vehicles (http://www.doi.org/10.17600/13030070). Main cruise results, geological context, details of fault rocks are provided elsewhere (Escartín et al., 2017; Bonnemains et al., 2017).

The exposed, corrugated detachment fault extends ~7 km in the spreading direction (East-West), and ~5.5 km perpendicular to the extension (North-South), respectively (Figure 1a; Olive et al., 2019). The microbathymetric corrugations (Figure 1b) have a relief of up to ~10-20 m, and along-extension lengths of a few hundred meters to a maximum of 2 km (Parnell-Turner et al., 2018). While the detachment fault surface is heavily blanketed by sediment and rubble, fault planes crop out on the flank of these corrugations, showing subhorizontal striations parallel to extension (Escartín et al., 2017). During the ODEMAR cruise, in situ fault rocks were sampled at seven outcrops distributed both along- and across-extension throughout the corrugated detachment fault surface (see numbered circles in Figure 1b). Among these outcrops, Outcrop #1 (Figure 1b) was in a ~70 m deep structural low within the detachment fault surface, which shows corrugations throughout. From these observations it was inferred that the fault zone is composed of anastomosing fault planes developing over a thickness of ~70 m or more (Bonnemains et al., 2017; Escartín et al., 2017; Parnell-Turner et al., 2018).

The activity of the Irinovskoe hydrothermal field was confirmed during the ODEMAR cruise (Escartín et al., 2017), ~1.8 km from the footwall cutoff (black circle in Figure 1b), at a location where samples of sulfides indicated recent or active hydrothermalism (Cherkashev et al., 2013; MacLeod et al., 2009; Pertsev et al., 2012). This site displays two black smoker vents at the summit of hydrothermal mounds (Active Pot and Pinnacle Ridge) venting fluids at ~365°C, in addition to several inactive mounds and chimneys (Escartín et al., 2017).

2.2. Detachment fault rocks: lithology and silicification

A total of 36 fault rocks were recovered in situ from the seven outcrops throughout the 13°20'N corrugated detachment fault using ROV Victor 6000, and are described in detail by Bonnemains et al. (2017). All the rocks are cataclastic breccias, with significant heterogeneity in clast abundance and size, and clast/matrix ratios, even at sample-scale (Figure 2). Most breccias contain solely mafic clasts (basalt/diabase; Figure 2a-d), and only 2 of the 36 samples display a mixed lithology, with coexisting mafic and ultramafic clasts (see Figures 1b and 2e-f). Several fault rock samples display highly localized deformation with striated slip plane surfaces (Figures 2c-e), as well as complex internal deformation textures, with cataclastic slip zones. Textures record, at the sample scale, several phases of both localized and distributed, penetrative deformation (Bonnemains et al., 2017).

Fault rocks are variably silicified indicating that (1) silica-rich fluids percolated through the fault rocks, and that (2) this silicification (and associated fluid flow) was likely heterogeneous at small spatial (outcrop) scale (Bonnemains et al., 2017). Moderately silicified mafic samples are clast-supported breccias, and contain clasts of hydrothermally altered mafic rocks with still identifiable magmatic textures. Most clasts display a relatively coarse doleritic texture underlined by fresh plagioclase laths (Figure 3b; see Bonnemains et al., 2017 for additional micrographs) indicating a diabase protolith. We have identified a single sample with a clast showing vesicular texture, corresponding to extrusive basalt (Figure 3a). Hence, the bulk of the fault material is incorporated from a dyke complex (Bonnemains et al., 2017), with limited reworking of shallow basalts, and consistent with an efficient exhumation (Olive et al., 2019).

The least silicified diabase clasts show rare relict clinopyroxene, largely replaced by amphibole of hornblende composition, associated to fresh plagioclase of labradorite composition, and no

associated chlorite (detailed mineral compositions in Bonnemains et al., 2017). This secondary mineral assemblage thus records hydrothermal recrystallization initiated under amphibolite facies conditions (≥ 500 °C). With increasing silicification, amphibole in clasts turns to actinolite, plagioclase becomes more albitic while chlorite crystallizes in clasts too, indicating greenschist facies conditions (~300°-500 °C). Clasts in these low to moderately silicified samples are surrounded by a matrix of finely crushed material, with newly formed chlorite and scattered quartz grains (<10 vol.% quartz). Highly silicified mafic samples are matrix-supported breccias with highly recrystallized clasts whose primary texture is largely obliterated. These breccia clasts are made up almost exclusively of chlorite and quartz, and are surrounded by a quartz-dominated (>90 vol.% quartz) matrix that also contains chlorite. Of the two samples bearing ultramafic clasts, only one has been silicificied. Abundant sulfides associated with quartz are found in three silicified fault rocks.

Cathodoluminescence imaging of selected samples also shows that quartz grains record successive fracturing and recrystallization episodes (Bonnemains et al., 2017), thus consistent with syntectonic quartz crystallization. Silicification and chloritization are closely associated during the alteration of the 13°20'N detachment fault mafic breccias. Indeed, chlorite is absent from the freshest diabase clasts (Bonnemains et al., 2017), chlorite content is low in moderately silicified samples and increases significantly with degree of silicification. Chlorite crystals imbricate quartz ones, or are included within quartz crystals, demonstrating co-crystallization in both clasts and matrix. Coeval quartz and chlorite crystallization is also unequivocal within late chlorite-filled veins crosscutting mafic samples. Thus, these microtextures record coeval formation of both minerals during detachment exhumation, and indicate that silicification occurred at greenschist-facies conditions (Bonnemains et al., 2017).

2.3. Samples selected for fluid inclusion and geochemical study

This study is based on a set of 6 representative detachment fault rocks that have been selected in order to (1) obtain a spatial coverage throughout the 13°20'N corrugated surface, and (2) cover various degrees of alteration and silicification (Table 1; Figure 1). These rocks were sampled in situ from five fault surface outcrops that are described in detail by Bonnemains et al. (2017; Figure 1b). All silicified samples contain abundant albeit very small fluid inclusions in quartz crystals.

Fluid inclusions suitable for this study were identified in 4 samples: three variably silicified breccias with only mafic clasts (ODM195, ODM155 and ODM218), and one silicified breccia with mixed mafic-ultramafic clasts (ODM173). We benchmarked geochemical analyses with two additional unsilicified samples of mafic and mixed mafic-ultramafic material (ODM115 and ODM217, respectively). Sample texture and mineralogy are summarized in Table 1. These samples are heterogeneous, showing a complex history with several deformation phases (Bonnemains et al., 2017), thus subsamples were also taken in some cases at varying distances from slip surfaces, as indicated below. Furthermore, to obtain representative whole-rock chemical analyses, larger subsamples were taken for three samples (i.e., ODM218, ODM173 and ODM217), labeled wr (whole rock) in Tables 2 and S1.

ODM115 is a basaltic clast from an unsilicified breccia (Figure 2a). It displays a typical pillow lava microtexture, with skeletal olivine microphenocrysts, radiating dendritic plagioclase, very fine-grained clinopyroxene crystals and vesicles (Figure 3a). Vesicles are filled with chlorite, veins with epidote, and the ground mass contains chlorite and pumpellyite. This quartz-free clast of extrusive basalt is considered as a reference for mafic material incorporated into the fault zone. It is the sole unsilicified mafic sample recovered from the fault outcrops.

ODM195 is a moderately silicified clast-supported breccia (Figure 2b) with clasts composed of either chlorite only, or actinolite + plagioclase \pm chlorite displaying relict doleritic textures (Figure 3b). The matrix contains crushed sub-millimetric clast fragments surrounded by newly formed chlorite and 5-10 vol.% of quartz (Figure 3b). Anhedral quartz crystals, typically \sim 100 μ m in length, are either scattered or in local aggregates in the matrix.

ODM155 and ODM218 are two highly silicified, matrix-supported fault breccias (Figures 2c-d). Both display a striated surface (slip plane) and an underlying well-developed slip zone (penetrative deformation). No primary mineral is preserved in any of the clasts, which contain either chlorite + titanite or chlorite + quartz (Figures 3c-d). Rare clasts preserve a primary doleritic texture (Figure 3c) despite pervasive alteration and mineral replacement, thus indicating a diabase protolith. Anhedral and subhedral quartz grains surrounded by interstitial chlorite represent >90 vol.% of the matrix. Both samples contain abundant sulfides, ODM155 containing only pyrite, whereas ODM218 contains pyrite with minor pyrrhotite and chalcopyrite.

Three subsamples were taken from ODM218, at varying distance from the slip surface, to investigate relationships between sample geochemistry and distance to slip planes, which may act as preferential fluid flow zones. ODM218a is located within 1 cm from the striated slip surface (Figure 2d), and contains only rare and submillimetric sulfide grains and clasts embedded in a quartz-rich matrix. When present, clasts are mostly silicified. ODM218b is ~4-5 cm away from the slip plane, and displays clasts either silicified or quartz-free up to 1 cm in size, in addition to disseminated quartz grains in the matrix. ODM218c is a fragment dislodged from the lower surface of the sample and away (~12 cm) from the slip plane, and contains quartz-free clasts surrounded by a silicified matrix with sulfides. In this set of samples, matrix quartz grain sizes increase with the distance to the slip plane, and thus ODM218c contains the largest matrix quartz grains.

ODM217 is a quartz-free, brecciated talc-amphibole schist displaying a slip surface on one side and several internal slip zones (Figure 2e). Clasts are either made up of talc and amphibole or chlorite ± titanite (Figure 3e). Matrix mineralogy varies among slip zones, and is made of talc, serpentine or chlorite (Figure 3e), with disseminated spinels. To account for sample heterogeneity, two subsamples were taken (Figure 2e). Subsample ODM217a is a chlorite-rich slip zone adjacent to a striated slip surface, whereas subsample ODM217b is a talc-rich zone a few cm away from this same slip surface. For whole-rock geochemical analyses, we thus consider the unsilicified breccia ODM217 as a reference, as it contains both ultramafic and mafic clasts.

ODM173 is a matrix-supported breccia (Figure 2f) containing both mafic and ultramafic clasts, made of talc or chlorite ± titanite (Figure 3f). The matrix is made of chlorite and talc, with variable amounts of quartz at the sample scale, and scattered spinel and sulfides (chalcopyrite + pyrite; Figure 3f). As for ODM217, we selected two sub-samples to investigate this heterogeneity. ODM173a displays few small clasts (< 1 mm) surrounded by a silicified matrix made of chlorite and talc associated with anhedral quartz crystals (<60 vol.% quartz, Figure 3f), whereas clasts in ODM173b are more abundant and larger (up to 1-2 cm), with only rare quartz grains and minor chlorite in the matrix. Sulfides are more abundant in ODM173b than in ODM173a.

3. Methods

To investigate the composition of fluids circulating through the detachment fault zone, and that are responsible for pervasive silicification, a study of fluid inclusions trapped in quartz crystals was

coupled to whole-rock and mineral geochemical analyses (major and trace element). Fluid inclusion analyses included microthermometry, Raman spectroscopy, and LA-ICP-MS. We also used SEM-cathodoluminescence to study the growth history of quartz crystals hosting fluid inclusions. Microthermometric measurements of fluid inclusions determined isochoric relationships that, coupled with geologically constrained pressure intervals, provided estimates of fluid inclusion entrapment temperatures. These temperatures were then compared to silicification temperatures inferred from the composition of chlorite co-crystallized with quartz, using the geothermometer of Bourdelle et al. (2013).

3.1. Whole-rock and mineral chemical analyses

Whole rock chemical analyses were performed at the SARM-CRPG (Nancy, France). Major elements were analyzed by inductively coupled plasma optical emission spectroscopy (ICP-OES) after fusion with LiBO₃ and dissolution in HNO₃. Trace elements were quantified by inductively coupled plasma mass spectrometry (ICP-MS) following the procedure described by Carignan et al. (2001). To account for sample heterogeneity, subsamples were analyzed for ODM218, ODM173 and ODM217 (see section 2.3).

Major element composition of chlorite was analyzed by EPMA (CAMECA SX-FIVE, CAMPARIS, Sorbonne Université, Paris, France). Analytical conditions were 15 kV-10 nA in WDS mode, for analysis of major elements. Fe₂O₃ (Fe), MnTiO₃ (Mn, Ti), diopside (Mg, Si), Cr₂O₃ (Cr), orthoclase (Al, K), anorthite (Ca) and albite (Na) were used as standards for elements in parentheses. Only chlorite analysis with oxide sum in the 86-89 wt.% range and Na₂O + K₂O + CaO < 1 wt.% were considered. Structural formulas were derived on a 14-oxygen basis. Chlorite analyses with Si \leq 4 atoms per formula unit, cation sum \leq 10.1 apfu and vacancies >0.01 pfu were considered. In situ trace element analyses were carried out on 22 chlorites at GeoRessources (Nancy, France) using LA-ICP-MS. Technical details can be found with results in Table S3.

3.2. Fluid inclusion analyses

Fluid inclusions analyses were performed on 100 µm thick double-polished sections. Successive generations of fluid inclusions trapped during quartz crystal growth were first characterized using

an Olympus BX-50 transmitted light microscope at IPGP. Inclusions either isolated or clustered (i.e., distant less than five times their length; Figure 4a) are considered as primary according to the criteria of Roedder (1984) and Van den Kerkhof and Hein (2001), whereas those located along trails crosscutting quartz grains are considered as secondary (Figure 4b). The location of the different generations of fluid inclusions within individual quartz crystals was compared to crystal growth history as revealed by SEM-cathodoluminescence observations (as in Boiron et al., 1992), conducted at ISTeP - Sorbonne Université (Paris, France) on a Scanning Electron Microscope Zeiss Supra 55 equipped with an EDS system and an Oxford Instruments cathodoluminescence system.

Microthermometric fluid inclusion measurements were carried out at ISTeP - Sorbonne Université under an optical microscope equipped with a Linkam THMSG 600 heating-freezing stage with temperatures ranging from -196 °C to +600 °C. The stage is controlled by a Linkam TMS 93 programmer via LinkSys software v.2.15.

Cycles of repeated homogenization and ice melting temperature measurements were conducted on 176 individual inclusions to measure the temperature of phase changes for the gas-to-liquid homogenization (temperature of homogenization, Th) and for the ice melting point (temperature of final ice melting, Tm_{ice}), respectively. Several cycles performed to test reproducibility show that ice melting and homogenization temperatures were reproducible within 0.1 °C (~0.2 wt.% eq. NaCl) and 1 °C, respectively. Inclusions yielding non-reproducible measurements systematically showed complex morphologies and were thus discarded, yielding 119 fluid inclusions for this study. We performed temperature cycles in the potential range of hydrate melting temperatures, following Raimbourg et al. (2014), but no gas hydrate was detected. Salinity was estimated from Tm_{ice} using the equation in Bodnar (1993), assuming that the entrapped fluid is a pure H₂O-NaCl solution. Isochoric P-T relationships followed by fluid inclusions were derived from Th and NaCl molality, following the equation of Zhang and Frantz (1987), suitable for our P-T and salinity range.

Gas content of fluid inclusions was analyzed with a Dilor-Labram Raman microspectrometer at GeoRessources (Nancy, France) on a representative set of 83 inclusions. As all studied inclusions are two-phase, we focused the laser on the gas bubble to determine gas proportions. In these dominantly aqueous inclusions, H₂O vapor is by far the dominant gaseous phase. The relative molar proportions of the remaining minor gases (H₂, CO₂, CH₄, H₂S) were calculated with accuracy better

than 5% (Pasteris et al., 1988) following the procedure described in Dubessy et al. (1989). This involves band area integration at wavenumbers of each gas and gas specific volume, i.e., Raman scattering cross-section for each gas and instrument efficiency at the specific wavenumbers (Frezzotti et al., 2012). To subtract the air signal for N₂, we conducted blank analyses in the quartz immediately adjacent to each inclusion.

Fluid inclusion chemical composition was analyzed on all inclusions larger than 8 µm (i.e., 24 inclusions) with a LA-ICP-MS at GeoRessources (Nancy, France; see technical details in Table S3). We analyzed Na, K, Li, Mg, Fe, Mn, Cr, Co, Ni, and calculated absolute concentrations and limits of detection following Leisen et al. (2012), using Na content derived from microthermometric Tm_{ice} measurements (and calculated salinity) as internal standard. However, due to the low signal intensity for most measured elements, it was impossible to calculate accurate fluid inclusion element concentrations; these fluid inclusions were very small, with fluid released after 1-2 laser shots, and contained low salinity fluids that are therefore very diluted in metallic elements. While Na is likely the most concentrated cation, it was systematically difficult to detect by LA-ICP-MS due to ionization problems.

4. Results

4.1. Chemistry

4.1.1. Whole rock analyses

Major element compositions and selected trace element contents for samples described in Section 2.3 are presented in Table 2 (see Table S1 for full trace element analysis). Chemical data are compared with published data of mafic and ultramafic rocks recovered at the 13°20'N OCC (Wilson et al., 2013), and at the 15°20'N Fracture Zone (Godard et al., 2008; Paulick et al., 2006), although such comparison must be done with care as 13°20'N fault rocks are breccias, and differ from those at other sites, as presented above.

The unsilicified basaltic fault rock clast fragment (ODM115) has a composition similar to that of MAR basalts. In silicified mafic samples, the concentration of alkalis, Ca and Al (CaO in Figure 5a; Na₂O and Al₂O₃ in Figures S1a-b) decreases sharply with increasing Si content, with almost complete depletion in the highly-silicified samples ODM155 and ODM218. Mg# for basaltic

ODM115 and moderately silicified diabase ODM195 are in the range of MAR basalt and diabase values (Figure 5b). In contrast, Mg# clearly decreases in highly silicified samples ODM155 and ODM218 (Mg# < 45). Mixed mafic-ultramafic breccias show compositions that are intermediate between MAR mafic and ultramafic compositions, resulting from mafic and ultramafic material mixed in variable amounts (Figures 5a-b and S1a-b). As in mafic samples, CaO, Na₂O and Al₂O₃ contents, as well as Mg#, are significantly lower in the silicified sample ODM173 compared to unsilicified ODM217 (Figures 5a-b and S1a-b). Their SiO₂ content is higher than that of ultramafic rocks for both the silicified ODM173 and the unsilicified ODM217, which is talc-rich.

Cr and Ni concentrations (Figure 5c) in most mafic samples are similar to those of MAR basalts and diabases, with subsample ODM218b showing slightly higher contents in both Cr and Ni. For the two samples with both mafic and ultramafic clasts, Cr and Ni contents are intermediate between those of MAR mafic and ultramafic compositions, reflecting again their mixed lithologies. The REE patterns (Figure S1c) for all mafic breccias are flat, as observed for MAR basalt and diabase patterns (Wilson et al., 2013). While the unsilicified basaltic clast ODM115 displays REE-normalized values consistent with those of MAR mafic rocks, REE concentrations progressively decrease as silicification increases, but without any pattern modification, likely due to dilution by quartz crystallization. The two mafic-ultramafic fault rocks display REE patterns between those of mafic and ultramafic signatures, reflecting their mixed lithologies (Figure S1c).

Hence, bulk rock analyses reflect significant heterogeneity of breccia samples, controlled by both the degree of silicification and the nature of the clasts. However, such analysis only reveals relative elemental enrichment or depletion. We combine these chemical analyses to textural and mineralogical observations to constrain effective mass transfer, and to determine if silicification resulted from significant silica enrichment only, or was associated to efficient leaching of other elements. Several arguments indicate massive silica influx rather than an important leaching: (1) dilution effect observed for most trace elements (Figure S1c), (2) preferential preservation of initial textures with well-defined clast borders, which would have been erased by massive leaching, (3) the fact that the most silicified breccias are matrix supported (while less silicified ones are clast-supported), and (4) the preferential crystallization of quartz in the matrix. However, calcium and alkalis (and part of the aluminum) were almost completely leached from silicified mafic and mixed mafic-ultramafic samples, and this important decrease with increasing silicification cannot be explained solely by passive depletion due to quartz crystallization (grey arrows, Figures 5a and

S1a-b). Almost complete alkali leaching is also supported by mineralogical composition of all mafic fault rocks from the seven outcrops (Bonnemains et al., 2017). Indeed, highly silicified samples contain mainly chlorite and quartz, lacking mineral phases able to host alkalis (i.e., no plagioclase or amphibole in highly silicified samples ODM218 and ODM155). The decrease in Mg# for silicified samples (ODM155, ODM218 and ODM173) compared to unsilicified ones likely reflects iron enrichment, as Mg# would be unaffected by quartz crystallization (Figure 5b). This is supported by the growth of iron-rich sulfides, essentially pyrite, as confirmed by the low Cu and Zn content (Table 2).

4.1.2. Chlorite composition

Microprobe analyses show that chlorites from both mafic and mixed mafic-ultramafic samples contain 40-50 wt.% of FeO+MgO, 30-40 wt.% SiO₂ and 15-25 wt.% Al₂O₃ (Figure 6a). There is no systematic difference in chlorite composition between clasts and matrix for each sample (Figure 6a). Chlorites in moderately silicified ODM195 show slightly lower FeO+MgO and higher SiO₂ contents than highly silicified ODM218, while chlorite composition of highly silicified ODM155 overlaps both samples.

Chlorites contain Si apfu in the range 2.7-3.5, Al apfu is 1.5-2.6 and Mg# between 43 and 85 (Table S2). While Si and Al contents are clearly anticorrelated for all chlorites, Si content and Mg# are roughly correlated in each sample, although more scattered (Figure 6b). The unsilicified basalt sample (ODM115) has chlorite compositions similar to those from MAR basalts (Alt et al., 1985; Gillis & Thompson, 1993; Humphris & Thompson, 1978). Chlorite composition in moderately silicified breccia (ODM195), in both clast and matrix, is comparable to that in diabases (amphibolite facies; Castelain et al., 2014; Escartín et al., 2003; Figure 6b). For highly silicified samples (ODM155 and ODM218), chlorites clearly have a lower Mg# compared to ODM195, with a wider range (about 80-40), although most of them remain comparable to oceanic basalt chlorites (Figure 6b). Increasing silicification is obviously associated with iron enrichment and silica depletion in chlorites, a trend also observed by Saccocia and Gillis (1995) and Delaney et al. (1987) in the MARK area, and by Castelain et al. (2014) in diabase chlorite-quartz veins at the footwall of the Atlantis Massif detachment (Figure 6b). Indeed, iron-rich chlorites have been described in oceanic hydrothermal breccias and quartz veins associated with the dyke complex or the dyke-lava

transition (Alt et al., 1985; Delaney et al., 1987; Honorez et al., 1998; Humphris et al., 1998; Mottl, 1983; Saccocia & Gillis, 1995).

Concerning mixed mafic-ultramafic breccias, chlorites from the unsilicified sample ODM217 show the highest Mg# at ~80. This is expected in rocks containing fragments of ultramafic rocks, and are mostly comparable to chlorites from other oceanic detachment talcschits and amphibolite schists (e.g., 15°45'N, Escartín et al., 2003; South Atlantis massif, Boschi et al., 2006; Figure 6b). Mg# for chlorites in the silicified mixed breccia (ODM173) presents a broader range and lower values (Mg#~50-80) than those of talcschists (Mg#~80, Figure 6b), similar to the decrease in Mg# and Si with increasing silicification observed for mafic samples.

Concerning trace elements, chlorites from mixed mafic-ultramafic fault rocks are generally enriched in Cr, Ni, Co, but depleted in Ti, V, Mn compared to chlorites from mafic breccias (Tables S2 and S3), in line with literature data (e.g., chlorites from diabases and talcschists, Boschi et al., 2006; Escartín et al., 2003). We note that a few chlorites from highly silicified mafic samples (ODM155 and ODM218) also show enrichments in Cr, Co, and slight enrichments in Ni (Tables S2 and S3).

4.1.3. Calculated temperature of chlorite formation

Several chlorite thermometers exist in the literature, based on empirical, semi-empirical, or thermodynamic approaches. Empirical thermometers cannot be used here as their application is restricted to the chlorite compositional space used for their calibration (Bourdelle & Cathelineau, 2015). Furthermore, their applicability is questionable as they only consider one substitution, do not take into account the bulk rock composition, and have no thermodynamic basis (Bourdelle & Cathelineau, 2015). Thermobarometric models (e.g., Vidal & Parra, 2000; Vidal et al., 2001, 2005, 2006; Walshe, 1986), widely used in metamorphic environments, lack of thermodynamic data for the Si-rich Al-free chlorite end-member (Figure 6c), therefore precluding their use for our samples. Indeed, chlorites formed in the fault breccias (Figures 6b-c) have a relatively high Si content (2.7-3.5 apfu) and some chlorites plot in the clinochlore-sudoite-Al-free chlorite field (Figure 6c). We thus used the semi-empirical geothermometer of Bourdelle et al. (2013), suitable for all compositions of chlorite in equilibrium with quartz. This choice is also comforted by the quartz

and chlorite co-crystallization in both matrix and clasts during silicification (see 2.2 and Bonnemains et al., 2017).

The geothermometer of Bourdelle et al. (2013) is specifically calibrated for low-temperature contexts (i.e., T < 350 °C), through a linear equation linking the chlorite + quartz equilibrium constant to the temperature of crystallization, taking into account cationic substitutions involving Si and R²⁺ contents. For higher-grade contexts, several thermodynamic parameters (as the non-ideal contribution of the site mixing) cannot be linearized, and a quadratic correction (Bourdelle et al., 2013) is proposed instead ("Tcorrected" in Table S2). While these "Tcorrected" results should be cautiously considered (Bourdelle et al., 2013), this involves a small portion of our analyses.

Temperatures obtained with this thermometer mostly span the 150-400 °C range (except for a few outliers which are not considered). We find median temperatures of ~250 °C for the moderately silicified samples ODM173 and ODM195, and ~275 and ~300 °C for the highly silicified samples ODM155 and ODM218 respectively (Figure 6d; Tables S2 and S4).

4.2. Fluid inclusion results

4.2.1. Distribution and morphology of fluid inclusions

Fluid inclusions are common in quartz from the $13^{\circ}20$ 'N detachment fault rocks. Those in quartz crystals from clasts in highly silicified breccias are bigger and better preserved than those in quartz grains within the matrix, which shows abundant decrepitated inclusions. All inclusions contain aqueous fluid that is two-phase (liquid-vapor; Figure 4) at room temperature, with a vapor to total volume ratio ranging from 0.1 to 0.5 (Figure S2). In quartz grains distant from slip planes (>1 cm), fluid inclusions are abundant and mostly range in size from 8 to 12 μ m (Figure 4a). The contour of primary fluid inclusions is spherical to elongated, sometimes approaching negative crystal shapes (Figures 4a, d, e, i). No sign of decrepitation (e.g., Roedder, 1984; Touret, 2001) is noticeable, except in samples within ~1 cm from a slip plane (sample ODM218a), in which inclusions are small (<5 μ m), very irregular in shape, and often empty and probably decrepitated (e.g., Roedder, 1984). Inclusions in this sub-sample ODM218a were therefore discarded for microthermometric measurements.

Small and elongated fluid inclusions, a few microns thick and a few tens of micron long, are regularly distributed, defining trails. Their elongation is often slightly oblique to the trail direction (Figures 4b, c, f, g, h), but their vapor ratio is similar to that of primary inclusions (Figure S2). These correspond to recrystallized microfractures, and thus postdate primary ones described above (e.g., Roedder, 1984). These trails do not show any distinct orientation, neither relative to each other nor at the sample scale (Figures 4f, g). Most of them are intra-grain trails, as they remain within the limits of quartz grains (Van den Kerkhof & Hein, 2001), with few occurrences crosscutting grain boundaries (Figures 4f, g). These trails are thus related to quartz fracturing between successive quartz crystallization episodes as revealed by SEM cathodoluminescence (Figures 7 and S3; Bonnemains et al., 2017). Therefore, this second set of inclusions can be classified as "pseudo-secondary", following the criteria of Roedder (1984).

4.2.2. Microthermometry

Ice melting temperature (Tm_{ice}) measurements indicate that all inclusions, primary and secondary, contain moderately saline aqueous fluids. Tm_{ice} ranges from -6°C to -1.5 °C (Figure S4), yielding equivalent salinities ranging from 2.1 to ~10 wt.% eq. NaCl. Most fluid inclusions studied here thus have a salinity higher than that of seawater (3.2 wt.% eq. NaCl, blue line in Figure 8a; Table 3) with salinity restricted to the 4–6 wt.% eq. NaCl range for all samples but ODM218, which displays a wider salinity range (2.1–10 wt.% eq. NaCl). We note that the salinity is higher for subsample ODM218b (from 6.3 to 10 wt.% eq. NaCl) than for subsample ODM218c (from 2.1 to 6.1 wt.% eq. NaCl; Figure 8a).

Highly silicified mafic breccias (ODM155, ODM218) present higher homogenization temperatures (180–350 °C) than moderately silicified ones, either mafic (ODM195) or mixed mafic-ultramafic (ODM173; 150–220 °C; Figure 8a; Table 3). The highly-silicified sample ODM155 shows a wide range of Th values, spreading over ~130 °C, whereas the Th for all other samples show more restricted ranges of up to 90 °C. We do not observe any systematic difference in Tm_{ice} and Th between primary and secondary fluid inclusions in any of the samples (Figure 8a).

Intra-grain variations of fluid inclusion Th and salinities are shown in Figure 8b, where sets of inclusions from different quartz grains are shown with different colors. The strong variation of Th and salinity observed respectively for ODM155 and subsamples OM218b and ODM218c (highly

silicified mafic breccias) is thus observed not only at the sample scale, but also at the quartz grain scale (Figure 8b). SEM-cathodoluminescence observations reveal core-to-rim variations in quartz luminescence (Figures 7 and S3) that record successive steps of quartz crystallization (Bonnemains et al., 2017). The location of fluid inclusions displaying scattered microthermometric values on cathodoluminescence images (Figure 7b, d and S3b, e) suggests that fluid inclusions were trapped in quartz of varying luminescence, and thus during different quartz grain crystallization phases, between successive quartz fracturing episodes.

4.2.3. Composition of fluid inclusions

Fluid inclusions from the 4 fault rock samples studied here are mostly aqueous (i.e., H₂O-NaCl), as shown by the absence of gas hydrate detection. Raman spectroscopy analyses of the vapor phase (i.e., gas bubble) of aqueous fluid inclusions detected minor amounts of H₂, CO₂, CH₄ and H₂S (in addition to H₂O) in the highly silicified mafic breccias (ODM155 and ODM218; Table 3; Figure 9). Conversely, only H₂O vapor was detected in the moderately silicified samples ODM173 and ODM195, although the high fluorescence observed during Raman analyzes for the latter could have hidden small gas signal.

In sample ODM155, only two out of 17 fluid inclusions (Table 3) were not purely aqueous and contained traces of CO₂. On the other hand, 15 out of 28 inclusions from subsample ODM218b, and the 21 analyzed inclusions of ODM218c, contained traces of gases (other than H₂O), and dominated by H₂ (up to 100 mol.% of the H₂-CO₂-CH₄-H₂S content in 18 inclusions) in addition to variable proportions of CO₂ (up to 80 mol.% of the H₂-CO₂-CH₄-H₂S content), CH₄ (up to 12 mol.% of the H₂-CO₂-CH₄-H₂S content) and H₂S (up to 12 mol.% of the H₂-CO₂-CH₄-H₂S content, in 4 inclusions from ODM218c; Figure 9a). All inclusions display very low H₂S/H₂ ratios (Figure 9b), similar to those measured in ultramafic-derived hydrothermal vent fluids (Fouquet et al., 2010), while mafic hosted vents rather show high H₂S/H₂ ratios. We do not observe any correlation between gas compositions and microthermometric data (Figure S5).

LA-ICP-MS analyses were performed on a set of 24 fluid inclusions, but their small size precludes calculating reliable concentrations (see 3.2). K +/- Na and Li were detected in 10 inclusions, and associated to the presence of Cr +/- Ni, Co, Mn, Fe, Mg in 7 of them. Despite these limitations due to inclusion sizes, inclusion fluids in the 13°20'N detachment fault rocks appear to

be very diluted fluids. They can thus be assimilated to fluids in the H₂O-NaCl system, and hence their salinity can be estimated from the ice melting temperature (Bodnar, 1993).

5. Interpretation and discussion

5.1. Pressure and temperature conditions of silicification

5.1.1. Pressure range deduced from the geological context

Silicification occurred in the detachment plane, affecting cataclastic breccias of mainly mafic composition. Preserved doleritic textures in moderately silicified breccias indicate that these diabase rocks were part of the base of Layer 2 (dyke complex) at the hangingwall, underlying the rift valley floor, and mechanically accreted into the fault zone (Bonnemains et al., 2017). Geophysical observations along slow-spreading ridges suggest that Layer 2 thickness may vary between 1 and 2 km (see discussion in Bonnemains et al., 2017 and references therein), while a recent seismic experiment in this study area reports a Layer 2A thickness on-axis of ~2 km (Simão et al., 2020). Assuming a rift valley floor depth of ~3000 m, and considering seawater and crustal rocks densities of 1025 and 3000 kg.m⁻³ respectively, the pressure ranges for the base of Layer 2 can be estimated to 600-890 bars for lithostatic pressure, and 400-500 bars for hydrostatic pressure. Therefore, silicification and coeval fluid entrapment in quartz crystals in the detachment fault zone necessarily took place between the base of Layer 2 (400-890 bars) and the seafloor (300 bars; Figure 10).

5.1.2. Temperature of fluid inclusion trapping during silicification

Fluid inclusion isochores derived from microthermometric data (equation of Zhang & Frantz, 1987) represent the P-T relationship along which fluid inclusions were trapped (Figure 10), while the measured homogenization temperature Th (Figure 8) indicates the rooting temperature of the isochore on the liquid-vapor curve (from Sourirajan & Kennedy, 1962). Intersection of fluid pressure ranges (see above) with isochores provides an estimate of the temperature range of silicification and subsequent fluid entrapment in quartz crystals (Figure 10). For samples ODM195 and ODM173, temperatures of fluid entrapment are in the 160-280 °C range, while for highly silicified samples ODM155 and ODM218, entrapment temperatures are in the 200-400 °C and 275-

425 °C ranges respectively (Figure 10). As these fluid inclusions were trapped at various stages of quartz syntectonic crystallization (Figures 7 and S3), fluid entrapment temperatures do represent the temperature interval over which silicification occurred.

Significant scatter in isochores for ODM155 reflect Th variability. Indeed, ODM155 (and to a lesser extent ODM195) exhibits a broad range of homogenization temperatures (~130 °C) at both sample- and grain-scale, associated to homogeneous salinities (4-6 wt.% eq. NaCl; Figure 8). This pattern is typical of post-entrapment deformation of fluid inclusions (Roedder, 1984), and is common in most hydrothermal systems (Delaney et al., 1987; Kelley et al., 1993; Petersen et al., 1998). This result suggests that some inclusion cavities (and therefore fluid density) underwent later re-equilibration during penetrative deformation phases (i.e., slip zones and planes in both samples), slightly modifying P-T isochoric relationships (Diamond et al., 2010; Tarantola et al., 2010; 2012; Vityk & Bodnar, 1995). In contrast, ODM218 shows a very restricted Th range, thus the isochore fan rather reflects its high variation in salinity (2-10 wt.% NaCl, Figure 8) and thus in fluid density.

5.1.3. Comparison with the temperature of chlorite formation

The formation temperature of chlorite, which coevally crystallized with quartz in both matrix and clasts during silicification (see 2.2 and Bonnemains et al., 2017), is calculated from chlorite composition (see 4.1.3; Figure 6d; Tables S2 and S4), and is thus an independent estimate of silicification temperature. The large range of chlorite crystallization temperatures (~150-400 °C, Figure 6d) reflects continuous crystallization with quartz during breccia infiltration by hydrothermal fluids and exhumation towards progressively lower P-T conditions in the detachment plane. The temperature range of chlorite formation is coherent with the estimated trapping temperatures for fluid inclusions, intersecting in all cases the isochores at realistic pressures (Figure 10). Interestingly, the chlorite temperature range intersects the isochores at higher pressures, corresponding to lithostatic fluid pressures, for the moderately silicified samples (ODM173 in particular), than for highly silicified samples (ODM218 in particular) for which the median chlorite temperatures intersect the isochores at hydrostatic fluid pressures instead. Near hydrostatic fluid pressures are coherent with the significant fluid amount required to explain pervasive silicification, thus associated with an open system. In contrast, moderately silicified samples submitted to more

restricted fluid circulation may instead correspond to a system only transiently open. Silicification is both static and linked to deformation episodes, as demonstrated by successive steps of quartz growth-hydrofracturing-overgrowth (Figures 7 and S3). This complex system witnessed both spatial and temporal fluid pressure variations, bounded by end-member lithostatic to hydrostatic pressures, and linked to variations in rock permeability and fault zone connectivity, likely modulated by both silica sealing and hydrofracturing episodes (Bonnemains et al., 2017).

We thus interpret this silicification as a long-lived, complex process along the detachment plane during exhumation of the hangingwall-derived breccias (Bonnemains et al., 2017), consistent with the large temperature ranges of both chlorite crystallization and fluid entrapment in quartz. We also infer that silicification occurred mostly in the temperature range $\sim 200-400\,^{\circ}\text{C}$, the highest temperatures corresponding to the highly silicified sample spanning the largest salinity variations (2-10 wt.% NaCl, ODM218).

5.2. Fluid compositions and potential fluid sources

5.2.1. Insights from fluid inclusion salinity: phase separation

Fluid inclusions contain an aqueous fluid with salinities generally higher than that of seawater. Fluid inclusions from most samples (ODM155, ODM195, ODM173) present restricted salinity ranges (4-6 wt.% eq. NaCl) corresponding to ~115 to 170% of seawater salinity (3.2 wt.% eq. NaCl). This range is comparable to ranges of salinity (and Th) measured in fluid inclusions (Figure 8a) in rocks associated with hydrothermal fields both of mafic (TAG: Petersen et al., 1998; MARK: Delaney et al., 1987; Saccocia & Gillis, 1995) and ultramafic nature (Rainbow, Logatchev, Ashadze, Semenov, Irinovskoe; Bortnikov et al., 2011, 2014, 2015; Simonov et al., 2015). Fluid inclusions from the highly silicified sample ODM218 display a large salinity range (2.1 to 10 wt.% eq. NaCl, Figure 8a). At the grain scale, core-to-rim salinity variations are associated to luminescence variations (Figures 7b, d, S3b, e and 8b) but do not show any systematic pattern, rather suggesting that the fault zone witnessed pulses of fluids with variable salinity during deformation.

Salinity in fluid inclusions from subsample ODM218b reaches up to 3 times seawater salinity (i.e., 10 wt.% equiv. NaCl, Figure 8a). Similar values are also reported for MARK silicified breccias (Delaney et al., 1987). In contrast, many fluid inclusions of subsample ODM218c cluster

at seawater-like salinities, while others in the same sample have salinities below that of seawater (2.1 wt.% eq. NaCl, Figure 8a), as reported from TAG (Petersen et al., 1998). At least three mechanisms have been proposed to explain salinities of fluid inclusions differing from that of seawater: hydration reactions, chloride retention in secondary minerals, and phase separation:

- (1) Hydration reactions consume water and, consequently, residual fluids may be enriched in dissolved elements. These reactions increase salinity modestly, with +15% for basalt-seawater equilibration at 350 °C (Wetzel & Shock, 2000), therefore clearly insufficient to explain the high salinities recorded in sample ODM218b. This process is only efficient at low water-rock ratio conditions and closed systems (Delaney et al., 1987). However, important fluid fluxes are required to attain the observed high levels of silicification (up to ~90 vol.% quartz) in these matrix-supported breccias (see 5.2.2). This precludes that hydration reactions alone may significantly change salinities to values consistent with those observed here (Figure 8a).
- (2) Chloride storage in transient phases (such as amphiboles, up to 4 wt.% chlorine; Vanko, 1988) can also modulate the salinity of the circulating hydrothermal fluids (Kelley & Robinson, 1990; Kelley et al., 1992). At 13°20'N OCC, hydrothermal amphiboles formed under amphibolite facies conditions (i.e., in clasts, pre-dating silicification) are progressively replaced by chlorite during silicification at lower temperatures (greenschist facies), and could therefore increase fluid salinity. However, the low chloride content of these amphiboles (<0.29 wt.%; Bonnemains et al., 2017) suggests that dissolution of these phases during silicification cannot account by itself for salinities of fluid inclusions within mafic fault breccias, which are three times higher than those of seawater.
- (3) The only efficient mechanism to generate both high and low salinity fluids is the formation of brine and vapor phases by supercritical phase separation of either seawater or magmatic fluids, as invoked for high-temperature hydrothermal systems (e.g., Alt et al., 2010; Castelain et al., 2014; Delaney et al., 1987; Kelley & Delaney 1987; Kelley et al., 1992; Vanko, 1988). At the 13°20'N detachment fault, the minimum temperature for supercritical phase separation at the seafloor (~3000 m, 300 bars, see 5.1) is ~400 °C (Figure 11), and would form high salinity brines and a vapor phase. However, we do not have any direct evidence for in-situ phase separation during silicification as we lack fluid inclusions with pure brines (up to ~40-50 wt.% eq. NaCl), as observed in diabases (Kelley & Delaney, 1987) or trondhjemite (Castelain et al., 2014; Figure 8a), or

inclusions with pure vapor phase (vapor to total volume of 0.1-0.5 in our study). This suggests that phase separation occurred earlier, at deeper levels. Vapor and brine were likely segregated and migrated upwards separately, similarly to the two-stage model proposed by Delaney et al. (1987) for the MARK hydrothermal field. The wide range of salinities (i.e., from lower- to three times higher-than-seawater) reported in 13°20'N silicified fault rocks may be accounted for by either rehomogenization of variable amounts of brines and vapor phases once released in the detachment fault (in the one-phase field, Figure 11), or their mixing with fluids of different salinity, prior to fluid entrapment in quartz.

Although we have no direct constraints, we hypothesize that phase separation may have occurred in the brecciation zone or its immediate vicinity. Indeed, most of the breccia clasts from 13°20'N fault zone are originally diabase, suggesting that they were initially part of the hangingwall dyke complex adjacent to the detachment fault (see 2.2 and 5.1 and Bonnemains et al., 2017). With a fully developed amphibolite facies paragenesis (hornblende + labradorite), these rocks experienced hydrothermal alteration at temperatures higher than 500 °C (see review in Alt, 1995), likely at the base of the upper crust (root of the dyke complex), therefore acting as a reaction zone. Fluid pressures in this reaction zone may be between hydrostatic and lithostatic, even close to hydrostatic for high fluid fluxes. Figure 11 shows that cold seawater infiltrating the hangingwall towards the base of the crust would undergo phase separation between 440 °C and 570 °C, depending on the fluid pressure gradient. It is thus plausible that phase separation occurred in the reaction zone where brecciation is inferred to take place (e.g., Bonnemains et al., 2017). Diabase brecciation along the detachment fault zone may even enhance phase separation, promoting sudden fluid pressure drops leading to the crossing of the liquid-vapor curve (Figure 11).

5.2.2. Insights from bulk-rock and chlorite compositions: a mafic-rock derived fluid

Textural and mineralogical observations, coupled to bulk-rock analyses, document that diabase breccias underwent variable silicification during exhumation within the detachment zone (see 4.1.1), reaching up to ~90 vol.% of quartz in matrix-supported breccias (Figure 5; Tables 2 and S1). Such degrees of silicification require both circulation of a silica-rich fluid and high fluid-rock ratios along the detachment fault zone. Indeed, experiments suggest that water-rock ratios >50 are required for basalt replacement by quartz-chlorite assemblages at 300 °C (Mottl, 1983). Elevated

fluid fluxes are also supported by the enhanced leaching of alkalis and calcium (Figures 5a and S1a-b) from diabase clasts with increasing silicification (Bach et al., 2013; Cann, 1969).

Chlorite crystallization in both clasts and matrix is associated to silicification, as showed by chlorite and quartz co-crystallization textures, increasing amount of chlorite with silicification, and similar chlorite composition in both clasts and matrix (Figure 6a). The decrease of bulk-rock and chlorite Mg# with increasing silicification (Figures 5b and 6b), correlated to pyrite crystallization in the highly silicified breccias, suggests interaction with iron-rich fluids. The association iron-rich chlorites, quartz and pyrite has been observed in silicified oceanic hydrothermal breccias associated with mafic-rock hosted vents such as TAG (Alt et al., 1985; Honnorez et al., 1998; Humphris et al., 1998) or at the MARK area (Delaney et al., 1987; Saccocia & Gillis, 1995). For Saccocia and Gillis (1995), these minerals result from interactions with high salinity, silica- and iron-rich fluids, depleted in H₂S. This is in agreement with the high salinity (Figure 8) and low H₂S content (only recorded in 4 fluid inclusions from ODM218, Figure 9a) in our fluid inclusions. Hence, almost all H₂S was likely stored into pyrite, leading to iron-enriched fluids from which Fe-rich chlorites crystallized (type I breccias from Saccocia & Gillis, 1995). A low H2S content in fluids is also consistent with the small amount of metallic trace elements recorded in our fluid inclusions (i.e., diluted fluids, see 4.2.3). Such fluid composition is characteristic of highly evolved hydrothermal fluids sampled in upflow zones of hydrothermal cells, and recording mafic rock-hosted reaction zones (Bach et al., 2013; Saccocia & Gillis, 1995).

The significant amounts of silica precipitated in detachment fault breccias also suggest the leaching of a deep mafic source. Indeed, high-temperature hydrothermal fluids from the reaction zone at the base of the mafic upper crust may be quartz-saturated (or close to) according to Wetzel and Shock (2000; ~20 mmolal of aqueous silica at 400°C for seawater-basalt equilibrium). This is in agreement with fluid composition from basalt-hosted hydrothermal vents at >300 °C (Fouquet et al., 2010; Schmidt et al., 2011; McDermott et al., 2018). Fluids circulating upwards along the detachment fault zone may witness both a significant temperature reduction (to 200-300 °C) and pressure drops, promoting fluid supersaturation and quartz precipitation. On the contrary, hydrothermal fluids equilibrated with peridotite should be largely undersaturated with respect to quartz (0.5 mmolal of aqueous silica at 400°, Wetzel & Shock, 2000), and vent fluids from ultramafic-hosted hydrothermal systems generally record 6-8 mmol/L of aqueous silica (Fouquet, 2010). Under these conditions, a temperature drop of 200 °C will not be sufficient to trigger quartz

crystallization. Hence, silica may only be provided by fluids reacting with mafic rocks. Moreover, the high amount of silica crystallized in the breccia matrix (up to 90 vol% of quartz) requires circulation of extremely Si-rich fluids: phase separation and brine formation in the reaction zone may have enhanced aqueous silica concentration, as quartz solubility is about one order of magnitude higher in NaCl-rich brines than in pure H₂O (Scheuermann et al., 2018; Schmidt et al., 2011; Steele-MacInnis et al., 2011), promoting efficient silica transport towards the upflow zone along the detachment fault.

Therefore, significant amounts of highly evolved hydrothermal fluids from a reaction zone at the base of the hangingwall dyke complex flowed up along the detachment fault zone, transporting leached aqueous silica. This syntectonic circulation resulted in intense and pervasive silicification of mafic detachment fault breccias (with crystallization of Fe-rich chlorite and pyrite).

5.2.3. Insights from fluid inclusion gas content: mix with an ultramafic-derived fluid

While most fluid inclusions are purely aqueous (H₂O-NaCl), some contain small amounts of H₂, CO₂, CH₄, and minor amounts of H₂S (i.e., ODM218b and ODM218c; Figure 9; Table 3) in addition to dominant H₂O (H₂O-NaCl-H₂±CO₂±CH₄±H₂S). Seawater phase separation cannot produce all these gases, and other fluid sources are required. Among these minor gases, H₂ is by far the most abundant in the inclusions (Figure 9; Table 3). H₂ is likely produced during serpentinization reactions by oxidation of the iron contained in olivine, reactions that preferentially occur between 200 and 350 °C (Martin & Fyfe, 1970; McCollom & Bach, 2009; Seyfried et al., 2007). At high temperatures (>400 °C), H₂ may also be produced in mafic rocks by pyroxene alteration (Allen & Seyfried, 2003; Foustoukos & Seyfried, 2005). CO₂ can be either linked to seawater, or to a magmatic source enriching nearby fluids. CH₄ likely results from CO₂-rich fluids reacting with H₂. This reaction may occur either during serpentinization, from hydrothermal circulation in ultramafic rocks enriching fluids in both hydrogen and methane (Berndt et al., 1996; Boulart et al., 2013; Charlou et al., 1998, 2002; Holm & Charlou, 2001; Monnin et al., 2014; Wetzel & Shock, 2000), in H₂-rich fluid inclusions hosted in footwall magmatic rocks (e.g., Klein et al., 2019; McDermott et al., 2015), or at a later time (e.g., during transport), but in any case, prior to fluid entrapment during silicification.

To discriminate the origin of fluids that circulated along the detachment fault zone, we compare the ratios of H₂, CO₂, CH₄, H₂S in our fluid inclusions to those analyzed in vent fluids (from MAR mainly). While few inclusions present a CO₂-dominant gas content comparable to basalt-hosted vent fluids, most of these inclusions show high H₂/CO₂ and H₂/CH₄ ratios (Figure 9a) and very low H₂S/H₂ ratios (Figure 9b), signatures typical of ultramafic-hosted vent fluids (e.g., Rainbow, Logatchev, Semenov, Irinovskoe; Destringeville et al., 2015; Fouquet et al., 2010; Schmidt et al., 2011), and of hydrothermal alteration of ultramafic rocks (i.e., serpentinization < 400 °C, Wetzel & Shock, 2000). However, fluids from the Piccard vents at the ultraslow spreading Cayman ridge show anomalously high H₂ contents and H₂/H₂S ratios, for mafic-rock-derived vents (McDermott et al., 2018). These fluids are interpreted as resulting from basalt alteration in a reaction zone at very high temperature (>500°C) and high fluid/rock ratios (McDermott et al., 2018; Scheuermann et al., 2018). This suggests that the high H₂/H₂S ratios analyzed in our fluid inclusions could derive from the hangingwall reaction zone. Lacking experimental data on basalt alteration at >500°C, reactions able to release such high H₂ amounts remain unknow. The reaction invoked by Scheuermann et al. (2020), of amphibole destabilization to magnetite and quartz, is clearly incompatible with our rocks, as hornblende is fresh in the least altered clasts (Bonnemains et al., 2017), and no magnetite (neither quartz) was observed. Moreover, H₂/H₂S ratios in Piccard vent fluids remain lower than those measured in most of our fluid inclusions (Figure 9b), although there is no reason for different partitioning of H₂ and H₂S between brines and vapor phases after phase separation (McDermott et al., 2018; Scheuermann et al., 2020). Therefore, H₂ production in the reaction zone seems to be possible, but may not account for the elevated H₂/H₂S ratios analyzed in our fluid inclusions, which are best explained by H₂ derived from ultramafic-rock serpentinization. Contribution of ultramafic-derived fluids is further supported by slight Cr and Ni enrichments of both sample ODM218b (Figure 5c) and some chlorites from highly silicified mafic breccias (Tables S2 and S3; Angiboust et al., 2014; Boutoux et al., 2014; Locatelli et al., 2019; Spandler et al., 2011).

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While the hangingwall sampled by the fault zone likely extends only to the base of the dyke complex, the footwall below the fault zone and deep sections of the hangingwall likely contain mantle peridotites that witnessed fluid circulation. This is demonstrated by the occurrence of mixed breccias containing altered ultramafic clasts (Bonnemains et al., 2017), and the outcropping of ultramafic rocks throughout the area (Escartin et al., 2017; MacLeod et al., 2011). During

exhumation, peridotites likely interacted with seawater-derived fluids, and progressively serpentinized. Fluids may have been subsequently channelized into and along the detachment fault zone. Mixing of such seawater-salinity fluids with silica-rich brines and vapor phases (prior to entrapment) could account for the wide salinity range observed in trapped fluids from the silicified breccias (Figure 8). This contribution of serpentinization fluids must remain limited nonetheless, as the absolute gas amounts trapped in fluid inclusions are extremely low. We also note that two thirds of the analyzed fluid inclusions from ODM218 have a gas signature very close to that analyzed for Irinovskoe vent fluids (Figure 9a), and that present-day fluid circulation within the footwall may record similar fluid-peridotite reactions at depth, even though these fluids do not exploit the detachment fault as a flow channel.

5.3. Detachment faulting, potential heat sources at the ridge axis and scenario for fluid pathways

Our results suggest that silicification was linked to flow along the detachment fault zone of silica-rich brines and minor vapor phases likely mainly formed in the hangingwall reaction zone (Figure 12a). Such a reaction zone at the base of the upper crust requires a heat source, such as a magma chamber at deeper levels and nearby the fault, or an along-axis dyke propagation (Bonnemains et al., 2017; MacLeod et al., 2009). The conceptual model in Figure 12a builds on the detachment fault zone structure suggested by Bonnemains et al. (2017) and Parnell-Turner et al. (2018). This model includes a) the incorporation of mafic clasts from the hangingwall upper crust into the detachment fault zone, from its base to the near-surface (brown stars in Figure 12), b) a thickening of the fault zone during exhumation, and c) an anastomosed fault zone mixing footwall-derived ultramafic material with dominant mafic material from the hangingwall. This model is also consistent with recent 3D seismic experiments revealing elevated seismic velocities at the footwall, both along the exposed detachment fault, and below its continuation eastwards below the rift valley floor (Simão et al., 2020). Elevated footwall seismic velocities are consistent with a lithosphere composed of lower crust and upper mantle rocks, juxtaposed to mafic rocks across the detachment (Simão et al., 2020).

As illustrated in Figure 12b, we propose that seawater percolates downwards through the hangingwall crust, reaching a reaction zone at temperatures >500 °C, at or near the axial zone.

High-temperature fluids circulating into the reaction zone may hydrothermally alter surrounding mafic rocks, while releasing aqueous silica and potentially small amounts of H₂S in the fluid phase (e.g., Bach et al., 2013; Wetzel & Shock, 2000). At these P-T conditions, phase separation occurs and forms high-salinity brines and low-salinity vapor phases. The small number of inclusions recording lower-than-seawater salinity (Figure 8a) suggests that only limited amounts of vapor reached and circulated within the detachment zone, while most of these low-density vapor phases may have migrated upwards through the hangingwall crust (Figures 11 and 12). Denser, less buoyant brines may be stored at depth in crustal porosity, within diabase units (Fontaine & Wilcock, 2006; Fontaine et al., 2007). Brines may be released into the detachment fault zone upon diabase brecciation, together with minor amounts of vapor phases, locally produced H₂S, and potentially CO₂ either transported by seawater or exsolved from the magma lens. Diabase brecciation may enhance phase separation by causing sudden and local fluid pressure drops (Figure 11). In turn, fluid phase separation may also promote local overpressures that may favor brecciation along the fault (André-Mayer et al., 2002; Bertelli & Baker, 2010), with possible links to seismicity along the detachment.

Varying gas content and wide salinity ranges of trapped fluids point to a complex and dynamic system. During silicification of the fault zone, upwelling silica-rich brines and minor vapors mixed in variable amounts with H₂(± CH₄)-rich fluids derived from serpentinization of ultramafic material from the footwall (Figures 11 and 12; Andreani et al., 2014; McCaig et al., 2010; McCaig & Harris, 2012). These fluids homogenized as a single fluid phase before trapping (one-phase fluid field, Figure 11). Fluid fluxes from these different sources were likely variable both in time and within the fault zone, resulting in strong variability of both gas content and salinity of fluid inclusions (Figures 8 and 9). Significant variations of inclusion fluid composition at quartz grain scale (Figures 7b, d, S3b, e and 8b) clearly indicate that the fluids circulating along the detachment were inherently heterogeneous at small spatial scales and short temporal scales. Yet there is no evidence for any systematic long-term evolution in fluid sources or fluid mixing processes during silicification (i.e., similarity in primary and secondary fluid inclusion salinity ranges). Moreover, the wide distribution of samples along axis and the pervasively silicified fault zone also indicate that this heterogeneous flow was maintained over long-periods of time, and was sampled by the detachment fault along most of its along-axis length.

All these fluids were preferentially channelized within the permeable fault zone, and silica precipitation was likely triggered by a combination of temperature decrease, pressure drops, and mixing of silica-rich brines with less-salted fluids. Both chlorite and fluid inclusion temperatures (Figures 10 and 11) are lower than those measured at the base of the dyke complex elsewhere (Hole 1256; Alt et al., 2010), documenting a significant temperature drop and silicification occurring under greenschist facies conditions. Upflow silica-rich brines were likely cooled either continuously or episodically, during transient circulation in a complex, anastomosing fault zone, and coeval with deformation leading to quartz fracturing-sealing episodes (i.e., fluid pressure varying between hydrostatic and lithostatic; Figures 7 and 10), but also by fluid mixing with cooler fluids from the footwall serpentinization and potentially seawater percolating from the seafloor.

5.4. Fluids trapped in detachment fault rocks vs. seafloor hydrothermal venting

Models of fluid flow and detachment faulting often assume a permeable fault zone efficiently channeling fluids to the seafloor or the shallow crust (e.g., Andersen et al., 2014; McCaig & Harris, 2012; Tao et al., 2020). In these models, fluid flow through the detachment fault feeds hydrothermal systems found both at the footwall (e.g., Logatchev; Andersen et al., 2014; Longqi: Tao et al., 2020) and hangingwall (TAG; McCaig & Harris, 2012). The heat source animating this circulation is often located in the upper mantle, at the base of the detachment microseismicity.

Our data and results from the 13°20'N detachment are not consistent with these deep-rooted hydrothermal systems along detachment fault zones. First, the present-day Irinovskoe hydrothermal site is located on the striated surface. As the detachment fault emerges at <20° at the hangingwall cutoff and the detachment is capped by a highly fractured and deformed fault zone (e.g., Escartín et al., 2017; Parnell-Turner et al., 2018), no plausible mechanism can channel fluids from the active detachment fault zone below the rift valley floor (below the hangingwall cutoff) to the Irinovskoe site, over a horizontal distance of ~2 km. The footwall also displays both a high-velocity seismic anomaly underlying Irinovskoe, restricted to the shallower 2 km, and a broad low velocity zone at 7-10 km depth (Simão et al., 2020). Owing to seismic resolution limitations, the presence of magma chambers or heat sources of sizes below the seismic resolution (1 to 3 km in size, depending on the depth) cannot be excluded. We propose that the fluid circulation system underlying Irinovskoe is unrelated to the active detachment, and instead is fully hosted within the footwall, probably reaching depths >2 km. While we lack direct seismic evidence here, this model

is similar to that of Rainbow, a hydrothermal field at the top of a detachment massif that is underlain by seismic reflectors identified as active and fossil magma chambers at depths >3 km (Canales et al., 2017; Dunn et al., 2017).

Silicified fault rocks are distributed throughout the detachment, displaying similar fluid compositions in quartz inclusions. This observation points to a flow organization along the detachment that has not been recognized before. Geological observations demonstrate that syntectonic silicification and associated fluid circulation occurred along the whole length (along-axis, Figure 1) of the active detachment, and that it operated continuously over time (across-axis; Bonnemains et al., 2017). Fluid inclusions further indicate that similar fluid sources (similar primary and secondary inclusions) mixed within the detachment plane (see 5.2.1 and Figure 8a). This type of distributed and pervasive hydrothermal circulation, extending laterally over several km and sustained over long periods of time, contrasts with the extremely localized outflow observed at the seafloor over relatively small surfaces (spatial scales of a few hundreds of m at most). Assuming that similar fluid circulation linked to deformation operates along the present-day, active detachment fault and at depth, this would require an extreme fluid flow localization on a short vertical distance (~2 km), in addition to implausible fluid transport within the footwall.

In our preferred interpretation, the fault zone and the fluids within cool down efficiently during exhumation at the shallower levels of the fault. This cooling can effectively suppress convection and any active circulation in shallow parts of the system, precluding discharge of high-temperature fluids at the seafloor. We suggest that the observed present-day hydrothermal activity observed on the surface of footwall of detachments (e.g., Rainbow, Semenov, Irinovskoe, Mount Den, Mount Fuji) is most likely due to heat sources underlying these sites and located within the detachment footwall, rather than fluids steered along the detachment fault and with significant lateral flow. The hydrothermal system that we document here may be indeed active at depth at 13°20'N, and be widespread at other systems. Indeed, many active detachments rooting near-axis (e.g., TAG, Dragon Horn) may be active in close proximity to nearby heat sources (melt lenses or dykes propagating along-axis), sampling fluids from the associated reaction zone at depth. As these systems may be restricted to deep lithospheric levels (>1 km) and lack a seafloor expression, they may only be identifiable studying in situ detachment fault rocks.

6. Conclusions

The 13°20'N detachment fault zone displays pervasive, syntectonic silicification of fault breccia that is originally mostly of mafic origin (diabase). Quartz crystals in breccia matrix and clasts preserve fluid inclusions. Their analyses, coupled to both bulk-rock and chlorite geochemical compositions, allow us to constrain hydrothermal fluid flow during deformation, and propose a conceptual model of detachment deformation and fluid flow.

First, we document the mixing of two fluid sources based on fluid inclusion salinity and gas content: 1) primarily brines and minor amounts of vapor phases resulting from phase separation at high temperature (T >410 °C), and likely occurring in the hangingwall reaction zone at the base of the dyke complex; and 2) minor amounts of fluids that interacted with ultramafic rocks, likely recording serpentinization reactions at temperatures of 200-350 °C, associated with H₂ (and CH₄) production. The lack of correlation between salinity and gas content, as well as quartz grain-scale variations of fluid composition, suggests syntectonic mixing in variable proportions of compositionally heterogeneous flows at small spatial and short temporal scales.

The hydrothermal circulation we document along the detachment fault zone reaches the base of the hangingwall dyke complex, i.e., 2 km depth, as constrained by seismic refraction data. Fault zone rocks are brecciated and incorporated into the detachment, together with the silica-rich brines emanating from a nearby reaction zone. A significant temperature drop is recorded by chlorite thermometry, from amphibolite facies (~500 °C) to quartz/chlorite equilibrium temperatures as low as ~200 °C. This temperature drop likely occurs both over small spatial distances and over short periods of time, with transients, promoting syntectonic silica precipitation in the fault zone. The hydrothermal circulation documented here is decoupled from present day hydrothermal activity at the seafloor. We suggest that detachment-related hydrothermal flow, which is widespread, is likely hosted within the footwall of detachments, passively cross-cuts inactive detachment faults, and is unrelated to the active detachment fault operating at depth.

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Tables

Table 1. Characteristics of the six fault rocks considered for this study. The outcrop numbers refer to Figure 1b and to Bonnemains et al. (2017).

	Sample #				Dont!	Doole					Analys	es
Outcrop	ODM_ROC_ Referred to as	IGSN	Latitude	Longitude	Depth (m)	Rock type	Texture	Mineralogical composition*	Quartz content	Whole rock	Chlorite	Fluid Inclusi
1	V547_115 ODM115	CNRS0000000115	13°18.96' N	-44°53.43' W	3159	mafic	clast of breccia with vacuoles	Chl, Pl, Px, Pmp	absent	X	X	
2	V552_217 ODM217	CNRS0000000221	13°19.22' N	-44°53.31' W	3113	mafic / ultramaf ic	matrix-supported breccia with a striated surface and 4 slip zones	clast: Tlc / Chl ± Ttn matrix: Tlc, Amp, Srp	absent	x	X	
	V552_218 ODM218	CNRS0000000222	13°19.22' N	-44°53.28' W	3133	mafic	matrix-supported breccia with a striated surface and 6 slip zones	clast: Chl ± Ttn / Chl + Qz ± Sulf (Py, Po, Ccp) matrix: Qz + Chl + Sulf (Py, Po, Ccp)	high	X	X	х
3	V551_173 ODM173	CNRS0000000176	13°20.40' N	-44°54.03' W	3218	mafic / ultramaf ic	matrix-supported breccia with one side mostly matrix and the other containing clasts	clast: Tlc, Amp / Chl ± Ttn matrix: Tlc, Chl ± Qz ± Sulf (Py, Po)	high on matrix- supported side, mo- derate on the other	х	Х	
4	V551_195 ODM195	CNRS0000000199	13°19.51' N	-44°53.75' W	2922	mafic	clast-supported breccia	clast: Pl + Amp + Chl matrix: fine-grained with sub- mm clasts and rare Chl and Qz grains	moderate	X	X	2
6	V550_155 ODM155	CNRS0000000157	13°19.40' N	-44°54.04' W	3104	mafic	matrix-supported breccia with a striated surface and 5 slip zones	clast: Chl ± Ttn / Chl + Qz ± Sulf (Py) matrix: Qz + Chl + Sulf (Py)	high	X	х	2

^{*} Mineral abbreviations: Amp, amphibole; Ccp, chalcopyrite; Chl, chlorite; Pl, plagioclase; Pmp, pumpelliyte; Px, pyroxene; Py, pyrite; Po, pyrrhotite; Qz, quartz; Srp, serpentine; Ttn, titanite; Sulf, sulfide; Tlc, talc

Table 2. Major and selected trace element whole-rock analyses of mafic and mixed mafic-ultramafic breccias from the detachment fault surface (see Table S1 for full analyses).

N	Vature	Basaltic clast	Mafic	Mafic	Mafic	Mafic	Mafic	Mixed	Mixed	Mixed	Mixed	Mixed	Mixed
Degree of	f silicification	Qz free	Moderately Si	Highly Si	Highly Si	Highly Si	Highly Si	Qz free	Qz free	Qz free	Moderately Si	Moderately Si	Moderately Si
S	ample	ODM115	ODM195	ODM218a	ODM218b	ODM218 wr	ODM155	ODM217a	ODM217b	ODM217 wr	ODM173a	ODM173b	ODM173 wr
SiO	2 wt.%	51.81	49.16	84.35	62.40	65.72	79.27	49.41	55.08	48.90	55.64	57.37	68.25
Al ₂ O) ₃ wt.%	13.76	13.05	2.69	4.12	3.78	2.43	5.46	2.26	5.25	9.05	1.50	5.18
Fe ₂ O)3 wt.%	8.16	10.76	6.30	19.97	17.09	9.47	11.25	8.07	12.50	13.79	10.04	11.07
Mn(O wt.%	0.16	0.22	0.06	0.08	0.08	0.06	0.33	0.39	0.45	0.11	0.07	0.07
Mg(O wt.%	6.08	11.88	2.43	4.33	4.08	2.10	22.70	22.93	21.30	11.24	20.49	8.18
CaC) wt.%	11.27	4.57	0.28	0.48	0.30	0.28	4.32	6.72	5.33	0.24	0.09	0.17
Na ₂ (O wt.%	3.78	1.23	< D.L.	0.11	0.07	0.05	0.22	0.20	0.24	0.29	0.28	0.17
K ₂ C	wt.%	0.09	0.04	< D.L.	< D.L.	< D.L.	0.02	0.03	0.03	0.04	0.04	0.03	0.03
TiO	2 wt.%	1.37	0.65	0.17	0.27	0.26	0.15	0.42	0.12	0.40	0.49	< D.L.	0.26
P ₂ O	5 wt.%	0.16	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	0.06	< D.L.	0.05	0.05	< D.L.	0.04
PF	wt.%	3.03	7.22	3.44	7.72	7.77	6.67	6.39	4.86	6.52	9.18	10.28	6.24
Tota	al wt.%	99.65	98.79	99.71	99.48	99.14	100.49	100.60	100.67	100.97	100.11	100.15	99.66
Mg:	#	59.61	68.62	43.26	30.05	32.11	30.51	80.00	84.91	77.15	61.74	80.18	59.41
FeO	wt.%	4.74	5.54	5.50	14.46	12.49	3.51	8.01	5.61	8.85	7.45	5.75	6.14
S	wt.%	0.04	0.04	1.04	7.68	6.21	4.75	0.02	0.03	0.02	0.66	0.69	1.98
Cr	ppm	308.50	349.30	293.70	590.40	525.60	303.30	1826.00	1895.00	2070.00	934.30	3319.00	1214.00
Cu	ppm	66.84	11.68	87.36	564.10	697.30	19.05	874.20	450.80	1540.00	504.20	1609.00	605.20
Ni	ppm	112.80	126.40	140.40	492.70	385.90	190.70	1080.00	1180.00	1064.00	429.50	1609.00	710.70
Zn	ppm	60.02	105.60	45.73	56.51	64.18	33.78	139.50	115.00	161.80	237.90	2947.00	379.70

<D.L.: lower than detection limit. Qz: quartz; Highly Si: highly silicified</p>

 Table 3. Microthermometric measurements in fluid inclusions and Raman spectroscopy results.

Sample	Quartz	Nb		Th (°C)		Salinity	(wt.% NaCl	eq.)	Gases other than H ₂ O in fluid inclusions, analyzed by Raman spectroscopy
	location		Range	Average	SD	Range	Average	SD	
ODM195	Matrix	19	149-218	182	16	4.3-5.9	5.2	0.5	12 FI studied – no gas detected
ODM155	All	24	183-322	263	25	3.5-5.9	4.8	0.3	
	- Matrix	14	197-286	258	24	4.3-5.9	4.9	0.4	17 FI studied – 2 with CO ₂ only
	- Clasts	10	183-322	271	23	3.5-5.3	4.7	0.3	
ODM218b	All	30	260-336	318	9	4.8-10.2	7.6	0.9	
	- Matrix	6	260-329	305	25	6.2-6.9	6.4	0.3	28 FI studied – 7 with H_2 only. 6 with H_2 + CO_2 . 2 with H_2 + CH_4
	- Clasts	24	310-336	321	5	4.8-10.2	7.9	0.8	
ODM218c	Matrix	36	258-348	317	12	2.4-6.2	4.3	0.8	21 FI studied – 11 with H ₂ only. 1 with H ₂ +CO ₂ . 3 with H ₂ +CH ₄ . 2 with H ₂ +CO ₂ +CH ₄ . 4 with H ₂ +CO ₂ +CH ₄ +H ₂ S
ODM173	Matrix	10	153-177	176	13	3.9-6.4	4.6	0.8	6 FI studied – no gas detected

Nb: number of fluid inclusions; SD: standard deviation

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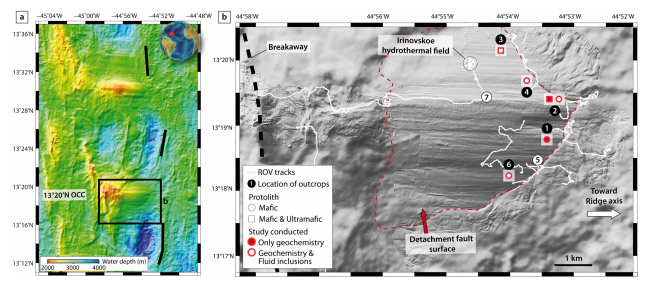


Figure 1: (a) Multibeam bathymetry data of the 13°20'N detachment region, showing the core complex (black box) and the ridge axis (thick black lines). (b) Shaded relief microbathymetry acquired with the AUV Abyss (GEOMAR), collected during the ODEMAR cruise (https://doi.org/10.17600/13030070) over the 13°20'N OCC, complemented with shipboard bathymetry. White lines are tracks of ten ROV dives. Fault surface outcrops identified and sampled with the ROV are shown by a circle (see Bonnemains et al., 2017). Samples used for this study come from five of them (black numbered circles and associated red symbols, see **Table 1** for details).

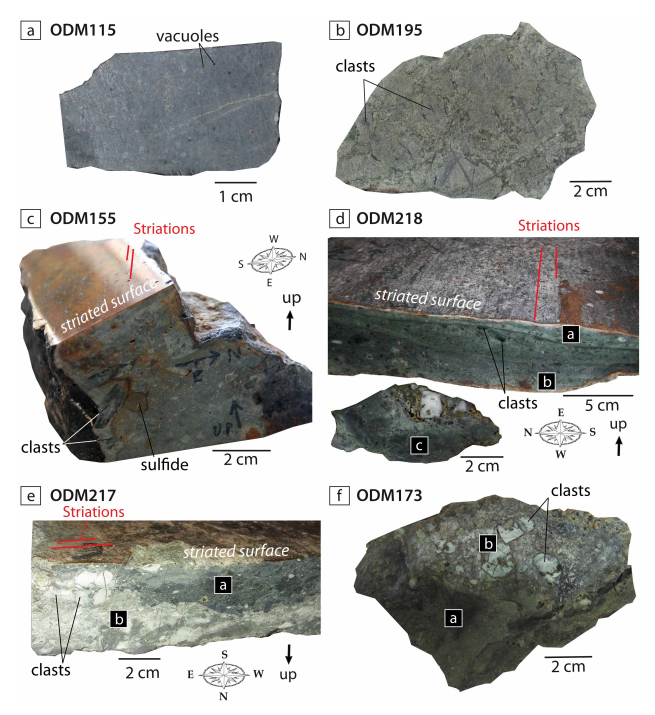


Figure 2: Macro-photographs of the studied fault rocks. (a) Clast of metabasalt with no quartz ODM115. (b) Moderately silicified mafic breccia ODM195. (c) Highly silicified mafic breccia ODM155. (d) Highly silicified mafic breccia ODM218 that is distinguished in two parts (subsamples a and b) at different distance from the striated surface (or slip plane) and a fragment dislodged from the lower part of sample (subsample ODM218c). (e) Unsilicified mafic-ultramafic breccia ODM217. We identify two parts based on the difference in color and mineralogical

composition between these slip layers. (f) Silicified mafic-ultramafic breccia ODM173 with ODM173a richer in quartz than ODM173b. All samples were recovered in situ at the detachment fault zone.

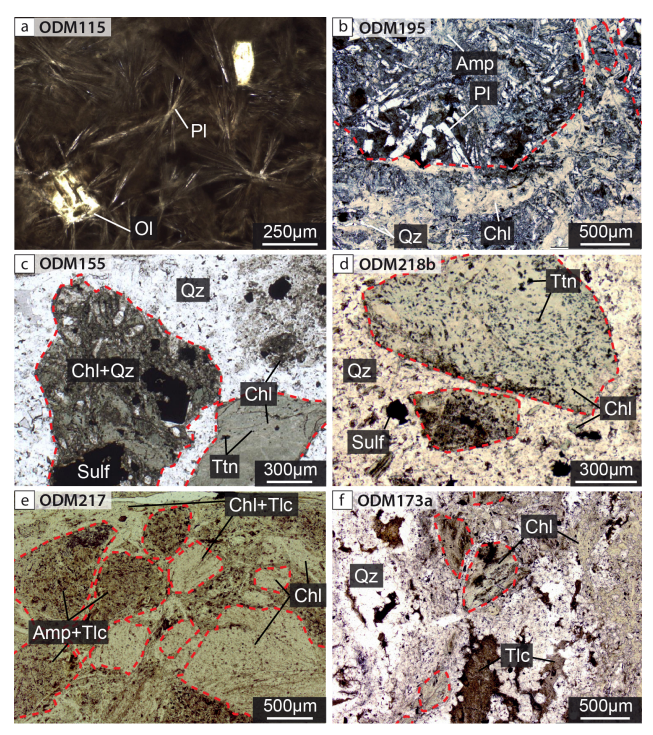


Figure 3: Microphotographs of studied samples in plane polarized light. (a) Unsilicified mafic clast showing basaltic texture with radiating plagioclase (Pl) laths and dendritic olivine (Ol) crystals. (b) Moderately silicified mafic breccia contains clasts displaying a doleritic texture, made of amphibole (Amp) + plagioclase ± chlorite (Chl), surrounded by a matrix consisting of crushed clasts mixed with chlorite and quartz (Qz). (c, d) Highly silicified mafic breccias containing clasts

of chlorite \pm titanite (Ttn) and chlorite \pm quartz \pm sulfide (Sulf), in a matrix mainly composed of quartz and minor chlorite and sulfide. In one of the clasts (c), the doleritic texture is still visible. (e) Unsilicified and (f) silicified mafic-ultramafic breccias containing both mafic (made of chlorite \pm titanite) and ultramafic (made of talc, Tlc, and amphibole) clasts embedded in a matrix composed of chlorite and talc, in addition to quartz for sample (f). (e) corresponds to zone a in Figure 2f. Clasts are delimited by red dotted lines.



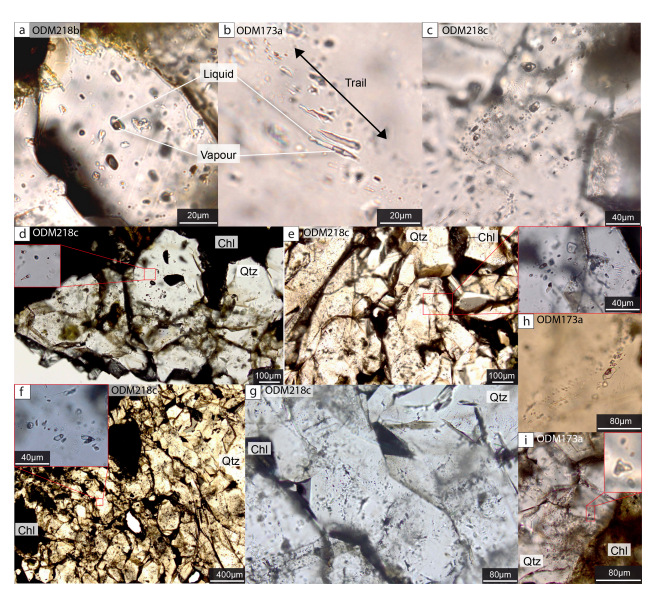


Figure 4: Microphotographs of representative two-phase (liquid and vapor) fluid inclusions in quartz grains. Clustered fluid inclusions with similar vapor/liquid ratios are considered as primary (a, d; e; i); fluid inclusions organized in trails are considered as secondary inclusions (b, c, f, g, h).

Note that most trails are intragrain trails, restricted to one quartz grain, while only few trails crosscut several quartz grain boundaries (g). Qtz: quartz; Chl: chlorite.

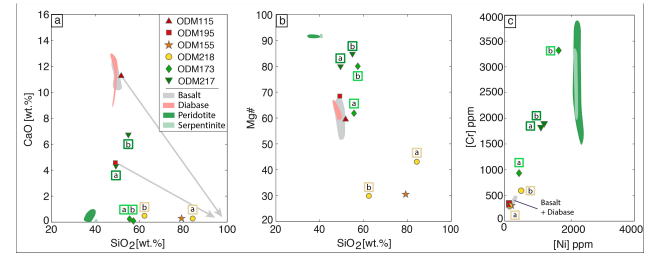


Figure 5: Bulk rock geochemistry. SiO₂ versus (a) CaO and (b) Mg# (=Mg/(Mg+Fe)*100); brecciated rocks show a relative decrease in both CaO and Mg# with increasing degree of silicification; mixed mafic-ultramafic breccias (green symbols) tend to be higher in magnesium at a given silica content; grey arrows: trend of passive depletion of CaO in breccias due to silica addition; Mg# will not be affected by silica addition. (c) Cr vs Ni; mixed breccias tend to be enriched in nickel and chromium compared with purely mafic breccias. Data for basalts and diabases from the 13°20'N OCC (Wilson et al., 2013) and for peridotites from the 15°20'N Fracture Zone (Godard et al., 2008; Paulick et al., 2006) are shown for comparison. Data are available in Table 2.

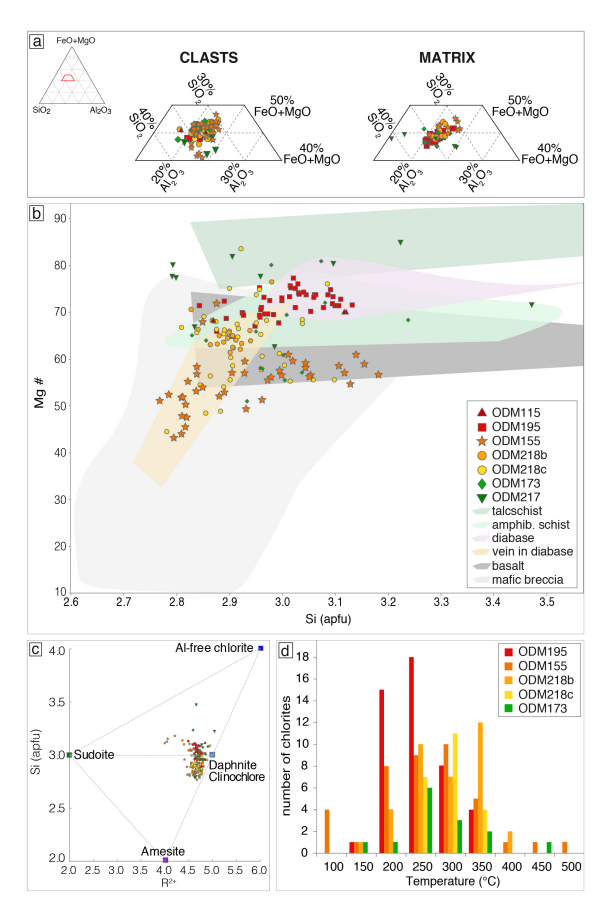


Figure 6: Chlorite composition in clasts and matrices (a) in a FeO+MgO, Al₂O₃, SiO₂ ternary diagram; (b) Mg# versus Si (atoms per formula unit); chlorite compositions are compared to other oceanic chlorites from ultramafic rocks (talcschists from 15°45'N: Escartín et al., 2003, and south of Atlantis Massif: Boschi et al., 2006); mixed mafic-ultramafic amphibolite schists (Escartín et al., 2003); mafic rocks (diabases: Escartín et al., 2003; basalts from MAR: Humphris & Thompson, 1978, and from MARK: Gillis and Thompson, 1993); altered mafic rocks (silicified breccias from MARK: Delaney et al., 1987; Saccocia and Gillis, 1995; chlorite-quartz vein in diabase, Atlantis Massif: Castelain et al., 2014); (c) Si versus R²⁺ diagram with chlorite endmembers; (d) Histogram of chlorite crystallization temperatures estimated using the semi-empirical geothermometer of Bourdelle et al. (2013). Chlorite analyses and calculated temperatures are available in Tables S2 and S4.

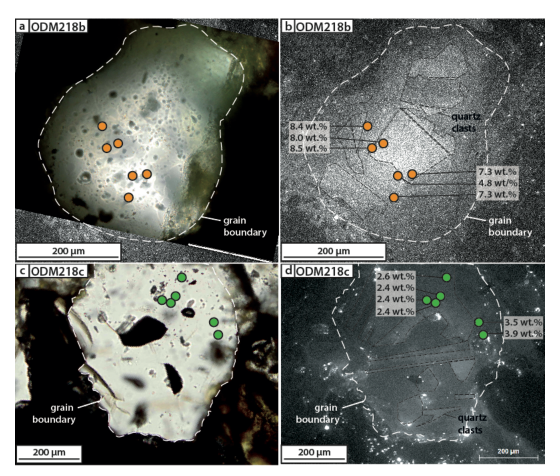


Figure 7: Distribution of measured fluid inclusions in quartz grains from ODM218b (a and b, orange dots) and ODM218c (c and d, green dots). Microphotographs under parallel nicols showing

individual quartz crystals in ODM218b (a) and in ODM218c (c). SEM-cathodoluminescence image of the same quartz grains in ODM218b (b) and in ODM218c (d); grain boundary shown by the white dashed line; luminescence variations illustrate that quartz grains are composed of multiple former quartz angular clasts (black dotted lines) around which quartz recrystallized in successive generations. Quartz crystallization was obviously syntectonic, with successive steps of quartz growth-hydrofracturing-overgrowth. The large salinity (indicated in wt.% NaCl equivalent) variation for fluid inclusions at the quartz grain scale suggests that fluids with different salinities were circulating (and thus trapped) during the successive episodes of quartz growth. Note that the position of fluid inclusions was projected on the grain surface, while inclusions are in fact distributed at various depths in the quartz grain.

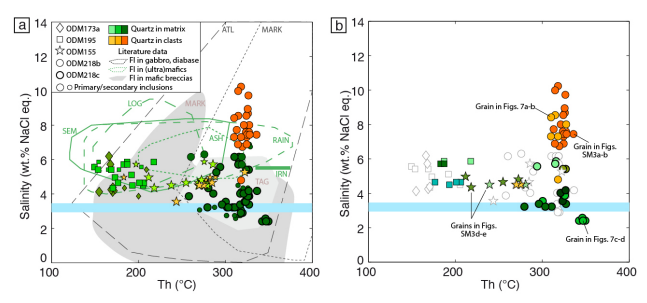


Figure 8: Homogenization temperatures (Th) against salinity for (a) all primary and secondary fluid inclusions; data are compared to Th and salinity ranges measured in fluid inclusions from oceanic gabbros (MARK: Kelley and Delaney, 1987), diabases (Atlantis Massif: Castelain et al., 2014), mafic silicified breccias from MARK (Delaney et al., 1987; Saccocia and Gillis, 1995) and TAG (Petersen et al., 1998), detachment plane (ultra)mafic rocks hosting hydrothermal vents (Rainbow, Logatchev, Ashadze, Semenov, Irinovskoe; Bortnikov et al., 2011, 2014, 2015; Simonov et al., 2015); (b) Heterogeneity of inclusion Th and salinity at the sample scale for the primary inclusions only, for clarity issues, using a similar color and symbol for inclusions in each grain. Seawater salinity is indicated with a blue line.

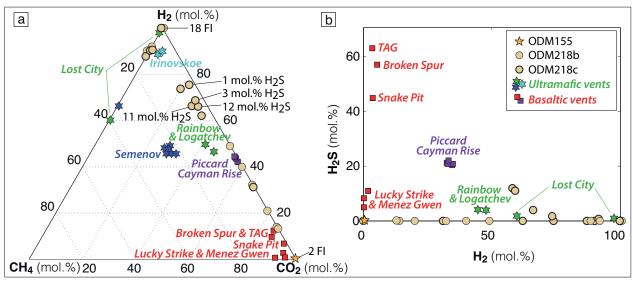


Figure 9: Molar proportion of gases (other than H₂O) analyzed by Raman spectroscopy in the vapor phase of two-phase fluid inclusions for the two highly silicified mafic breccias ODM155 and ODM218. All fluid inclusions are dominantly aqueous, and H₂O vapor is the dominant gas phase. (a) Relative molar proportions of CH₄, CO₂ and H₂ in the gas phase. Gases other than H₂O were analyzed in only 2 inclusions (over 17 analyzed) for ODM155 (and contain only CO₂ other than H₂O vapor), in 15 inclusions over 28 analyzed for ODM218b, and in the 21 fluid inclusions analyzed in ODM218c. 7 and 11 fluid inclusions contain only H₂ (other than H₂O vapor) in ODM218b and ODM218c, respectively. (b) H₂ versus H₂S content in fluid inclusions (expressed as a mol.% of CH₄-CO₂-H₂-H₂S in the vapor phase). Fluid inclusion gas compositions are compared to data from basaltic- (Menez Gwen, Broken Spur, TAG, Snake Pit, Lucky Strike) and ultramafic- (Rainbow, Logatchev, Lost City) derived hydrothermal vents (Fouquet et al., 2010). Data from Semenov and Irinovskoe ultramafic-derived vents (Destrigneville et al., 2015) and from Piccard mafic-derived vents on Cayman Rise (McDermott et al., 2018) are also plotted for comparison.

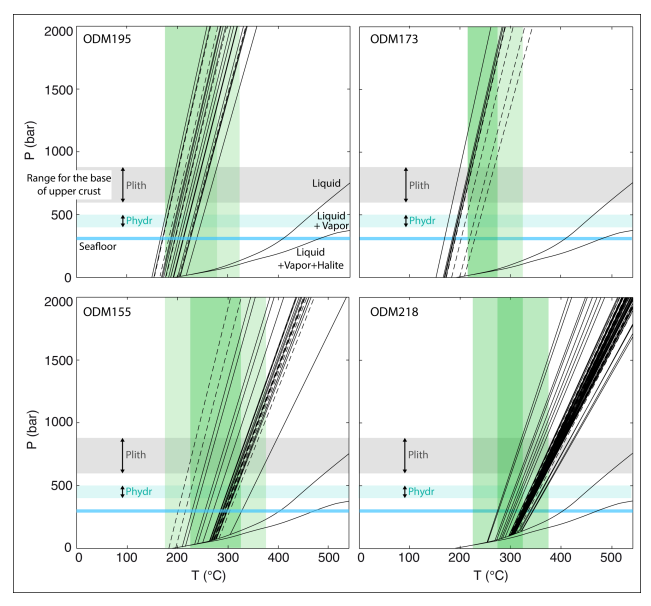


Figure 10: Range of P-T conditions for silicification of brecciated rocks within the detachment plane. P-T isochoric relationships for fluid inclusions, calculated from microthermometric data (equation of Zhang & Frantz, 1987), are plotted in black plain and dashed lines for primary and secondary fluid inclusions respectively. The liquid-vapor and liquid-vapor-halite curves are from Sourirajan and Kennedy (1962). Pressure at the seafloor is indicated with a blue line, and the range of lithostatic and hydrostatic fluid pressures at the base of the hangingwall upper crust (1–2 km thick) is in grey and blue respectively (assuming seawater and rock densities of 1025 and 3000 kg/m³ respectively). The temperature range of chlorite crystallization during silicification, calculated from chlorite composition (geothermometer of Bourdelle et al., 2013), is in green (darker green for higher number of chlorites; see Figure 6d).

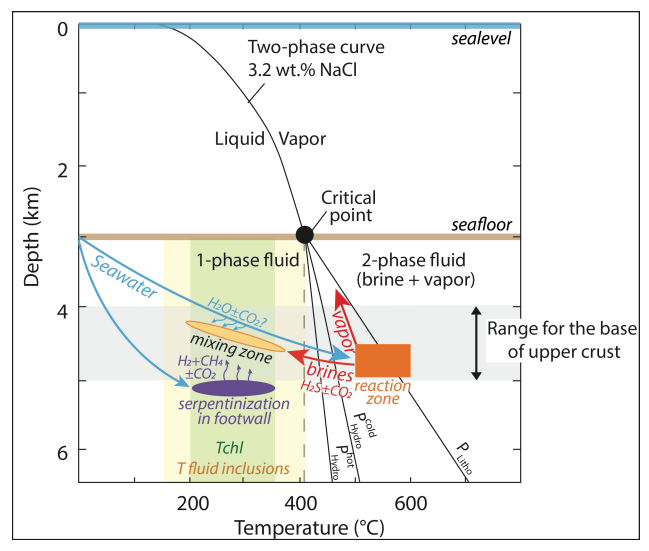


Figure 11: P-T-depth relationships in the NaCl-H₂O system for a hydrothermal seawater-like solution (3.2 wt.% NaCl). The two-phase curve separates the one-phase liquid field from the two-phase liquid+vapor field (Sourirajan & Kennedy, 1962). Temperature-depth relationship for the two-phase curve beyond the critical point (407°C, 298 bars, black dot) is calculated for both lithostatic and hydrostatic fluid pressures. Cold and hot hydrostatic pressure gradients (100 and 50 bar/km respectively) were taken from Coumou et al. (2009) and Castelain et al. (2014). Seawater (blue arrow) infiltrating the hangingwall crust (1-2 km thick) may undergo phase separation, at least when reaching the reaction zone (orange rectangle) at the base of the crust (diabase clasts suggest temperature >500 °C). While part of vapor phases can migrate upwards into the crust, higher density brines are likely released in the detachment plane upon diabase brecciation (red arrows). They are mixed in variable amounts with fluids derived from footwall serpentinization

(purple arrows) and potentially small amounts of seawater circulating in hangingwall basalts (blue arrows). Temperature ranges of chlorite formation (light green zone) and homogenization of fluid inclusions (light yellow zone) are reported (see Figures 6d, 8, 10 and text for details).

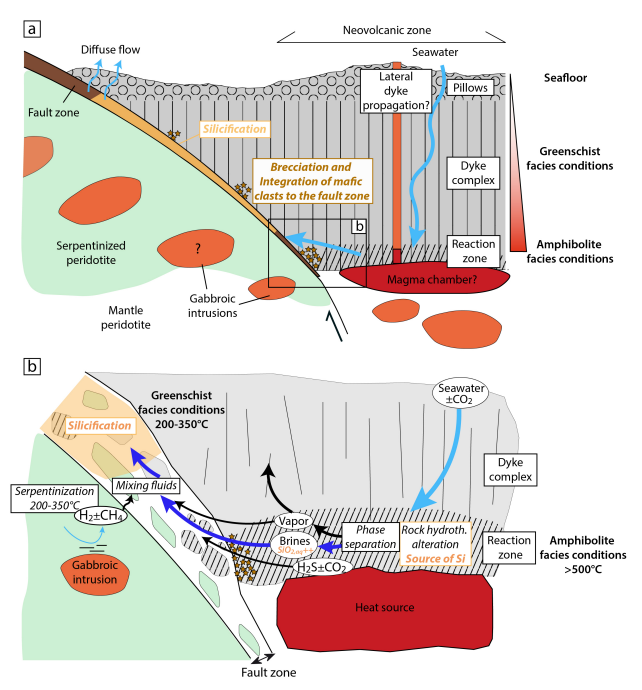


Figure 12: Schematic interpretation of fluid circulation along the 13°20'N OCC. (a) The hangingwall corresponds to a section of upper crust while the footwall progressively exhumes

material from deeper levels of the lithosphere (mantle-derived peridotites with gabbroic intrusions). A reaction zone at the base of the upper crust is generated by a heat source located beneath the neovolcanic zone. The fault zone thickens during exhumation, due to the integration of hangingwall material. (b) Close up of the reaction zone close to the heat source. Seawater percolates down to the reaction zone where increased pressure and temperature generate phase separation into brine and vapor phases. Brines are enriched in silica released by hydrothermal alteration of the mafic rocks. While most of the vapor phases escape towards the surface, brines (and a small portion of vapor phases) are integrated into the fault zone during the overplating and mix with (small amounts of) hydrogen-bearing fluids generated by serpentinization of the footwall. Reaction of hydrogen with CO₂ either dissolved in seawater or released by magmatic activity results in the formation of methane. As the fluid ascents and cools down, the solubility of silica strongly decreases resulting in precipitation of quartz that entraps the fluid inclusions.

@AGU PUBLICATIONS

1532	
1533	Geochemistry, Geophysics, Geosystems
1534	Supporting Information for
1535	Fluid circulation along an oceanic detachment fault: insights from fluid
1536	inclusions in silicified brecciated fault rocks (Mid-Atlantic Ridge at 13°20'N)
1537 1538	A. Verlaguet ^{1*} , D. Bonnemains ² , C. Mével ² , J. Escartín ^{2,3} , M. Andreani ⁴ , F. Bourdelle ⁵ , M-C. Boiron ⁶ , V. Chavagnac ⁷
1539 1540	¹ Sorbonne Université, CNRS-INSU, Institut des Sciences de la Terre, ISTeP UMR 7193, F 75005 Paris, France
1541	² Université de Paris, Institut de Physique du Globe, CNRS, F-75005 Paris, France
1542 1543	³ Laboratoire de Géologie (CNRS UMR 8538), Département de Géosciences, Ecole Normale Supérieure, PSL University, F-75005 Paris, France
1544	⁴ Laboratoire de Géologie LGL-TPE, Université de Lyon, UCBL, ENSL, CNRS, Villeurbanne, 69622, France
1545	⁵ Univ. Lille, IMT Lille Douai, Univ. Artois, Yncrea Hauts-de-France, ULR 4515 - LGCgE,
1546	Laboratoire de Génie Civil et géo-Environnement, F-59000 Lille, France
1547	⁶ Université de Lorraine, CNRS, GeoRessources, F-54000 Nancy, France
1548	⁷ Géosciences Environnement Toulouse (GET), Université Paul Sabatier Toulouse 3, CNRS UMR 5563, IRD,
1549	Toulouse, France
1550	* Corresponding author: anne.verlaguet@sorbonne-universite.fr
1551	
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1553	Contents of this file
1554	Figures S1 to S5
1555	Tables S1 to S4

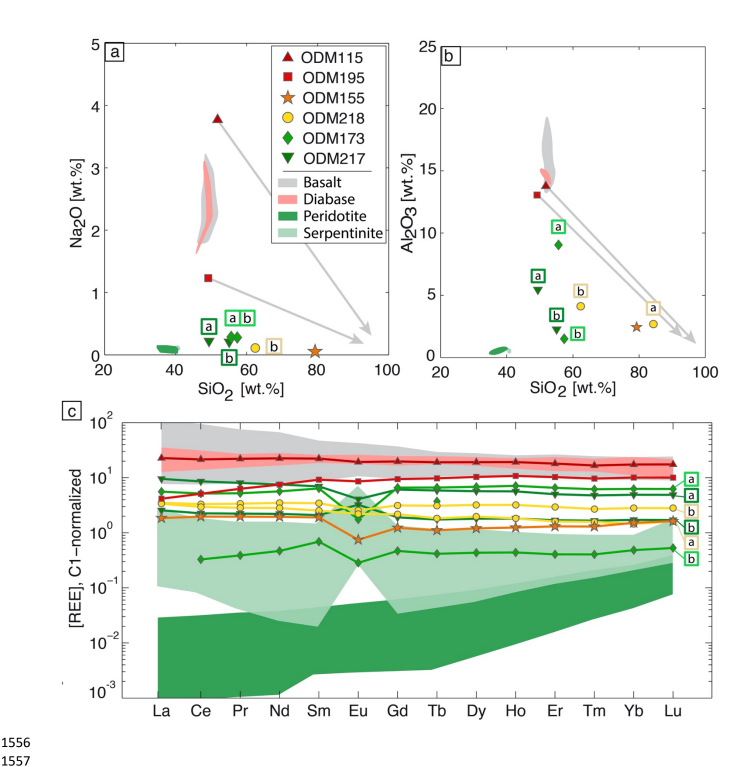


Figure S1: Bulk rock geochemistry. SiO₂ versus (a) Na₂O and (b) Al₂O₃; brecciated rocks show a relative decrease in both Na₂O and Al₂O₃ with increasing degree of silicification; grey arrows: trend of passive depletion of Na₂O and Al₂O₃ in breccias due to silica addition. (c) Rare earth element (REE) content of brecciated rocks normalized to chondrite. Compositions are compared to data for basalts and diabases from the 13°20'N OCC (Wilson et al., 2013) and for peridotites from the 15°20'N Fracture Zone (Godard et al., 2008; Paulick et al., 2006).

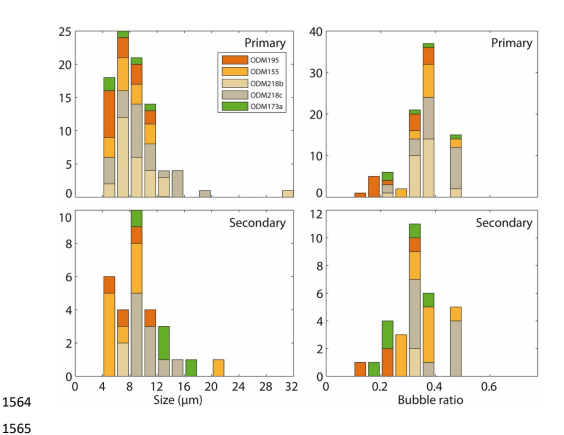


Figure S2: Size distribution (left) and bubble/inclusion volumetric ratio (right) histograms for primary (top) and secondary (bottom) fluid inclusions. Vertical axis is frequency.

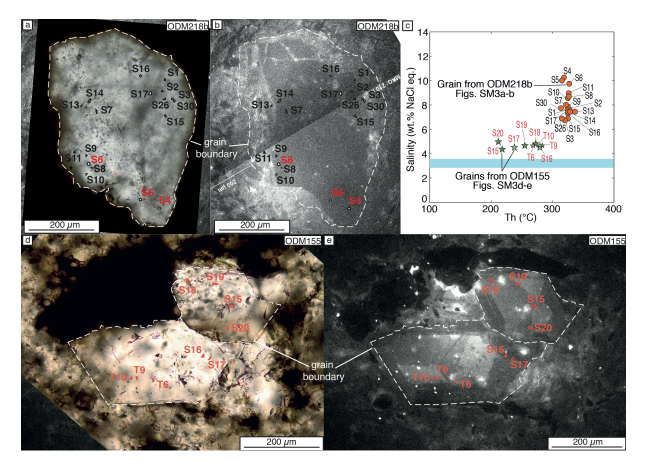


Figure S3: Location of analyzed fluid inclusions in quartz grains from ODM218b (a and b, orange dots in c) and ODM155 (d and e, green stars in c). Microphotographs under parallel nicols showing individual quartz crystals in ODM218b (a) and in ODM155 (d). SEM-cathodoluminescence images of the same quartz grains in ODM218b (b) and in ODM155 (e); grain boundaries are shown by the white dashed line; luminescence variations illustrate that quartz grains are composed of multiple former quartz angular clasts around which quartz recrystallized in successive generations. Quartz crystallization was unequivocally syntectonic, with successive steps of quartz growth-hydrofracturing-overgrowth. (c) Homogenization temperatures (Th) against salinity (indicated in wt.% NaCl equivalent) for primary (big symbols) and secondary (small symbols) fluid inclusions from quartz grains in ODM218b (a, b) and ODM155 (d, e). Seawater salinity is indicated with a blue line. The large salinity variation measured from fluid inclusions within individual grains indicates entrapment of fluids with varying salinities during successive quartz growth episodes. Note that the position of fluid inclusions was projected on the grain surface, while inclusions are in fact distributed at various depths within the quartz grain. In ODM218b (a, b) inclusions in red are those very close to the thick section surface.



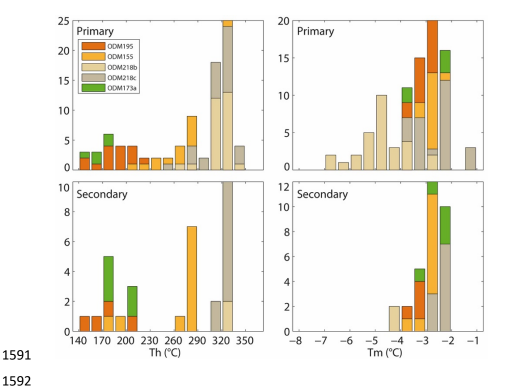


Figure S4: Histogram showing the distribution of homogenization temperatures (Th, left) and ice melting temperatures (Tm_{ice}, right) for primary (top) and secondary (bottom) fluid inclusions. Vertical axis is frequency.

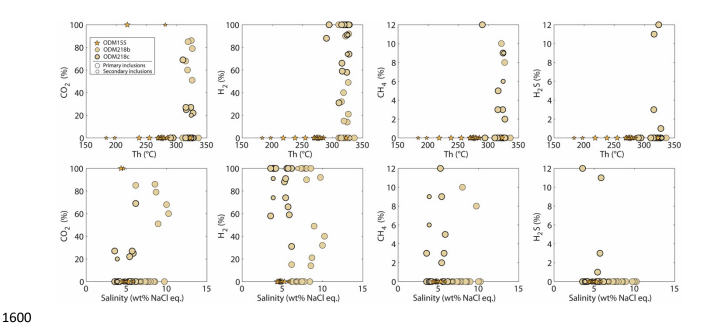


Figure S5: Gas content (mol.% of gas other than H_2O in the gas bubble) versus homogenization temperature (Th) and salinity.

Table S1. Full chemical analyses of the selected samples.

Nature		Basaltic clast	Mafic	Mafic	Mafic	Mafic	Mafic	Mixed	Mixed	Mixed	Mixed	Mixed	Mixed
Degree of silic		Qz free	Moderately Si	Highly Si	Highly Si	Highly Si	Highly Si	Qz free	Qz free	Qz free	Moderately Si	Moderately Si	Moderately Si
Sample		ODM115	ODM195	ODM218a	ODM218b	ODM218 wr	ODM155	ODM217a	ODM217b	ODM217 wr	ODM173a	ODM173b	ODM173 wr
SiO ₂	wt.%	51.81	49.16	84.35	62.40	65.72	79.27	49.41	55.08	48.90	55.64	57.37	68.25
Al_2O_3	wt.%	13.76	13.05	2.69	4.12	3.78	2.43	5.46	2.26	5.25	9.05	1.50	5.18
Fe ₂ O₃	wt.%	8.16	10.76	6.30	19.97	17.09	9.47	11.25	8.07	12.50	13.79	10.04	11.07
MnO	wt.%	0.16	0.22	0.06	0.08	0.08	0.06	0.33	0.39	0.45	0.11	0.07	0.07
MgO	wt.%	6.08	11.88	2.43	4.33	4.08	2.10	22.70	22.93	21.30	11.24	20.49	8.18
CaO	wt.%	11.27	4.57	0.28	0.48	0.30	0.28	4.32	6.72	5.33	0.24	0.09	0.17
Na₂O	wt.%	3.78	1.23	< D.L.	0.11	0.07	0.05	0.22	0.20	0.24	0.29	0.28	0.17
K₂O	wt.%	0.09	0.04	< D.L.	< D.L.	< D.L.	0.02	0.03	0.03	0.04	0.04	0.03	0.03
TiO ₂	wt.%	1.37	0.65	0.17	0.27	0.26	0.15	0.42	0.12	0.40	0.49	< D.L.	0.26
P ₂ O ₅	wt.%	0.16	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	0.06	< D.L.	0.05	0.05	< D.L.	0.04
PF	wt.%	3.03	7.22	3.44	7.72	7.77	6.67	6.39	4.86	6.52	9.18	10.28	6.24
Total	wt.%	99.65	98.79	99.71	99.48	99.14	100.49	100.60	100.67	100.97	100.11	100.15	99.66
Mg#		59.61	68.62	43.26	30.05	32.11	30.51	80.00	84.91	77.15	61.74	80.18	59.41
FeO	wt.%	4.74	5.54	5.50	14.46	12.49	3.51	8.01	5.61	8.85	7.45	5.75	6.14
H ₂ O total	wt.%	3.22	8.13	2.49	4.20	4.32	2.62	7.05	5.55	6.95	8.93	< D.L.	5.32
S total	wt.%	0.04	0.04	1.04	7.68	6.21	4.75	0.02	0.03	0.02	0.66	0.69	1.98
В	ppm	5	2	2	<2	<2	4	11	8	14	8	10	7
Cl	ppm	230	800	240	355	295	235	435	380	580	1 520	2 540	1 030
Li	ppm	3.7	7.6	5.4	3.0	3.3	8.3	7.4	7.6	7.4	4.3	4.2	3.1
As	ppm	2.29	< D.L.	< D.L.	< D.L.	< D.L.	1.80	1.88	1.75	3.25	1.68	< D.L.	< D.L.
Ва	ppm	35.37	4.17	3.00	1.67	< D.L.	2.66	< D.L.	1.69	< D.L.	1.90	< D.L.	6.75
Be	ppm	0.64	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	0.46	0.60	0.50	< D.L.	< D.L.	< D.L.
Bi	ppm	0.19	< D.L.	0.21	0.69	1.30	0.32	< D.L.	< D.L.	0.10	0.12	0.52	0.20
Cd	ppm	0.14	< D.L.	< D.L.	< D.L.	0.12	< D.L.	< D.L.	< D.L.	< D.L.	0.68	203.80	12.18
Ce	ppm	13.19	3.13	1.81	2.03	1.98	1.20	5.26	1.38	3.77	3.17	0.20	1.35
Co	ppm	30.87	31.10	36.76	206.20	151.30	82.21	71.51	61.76	71.78	49.51	280.50	60.67
Cr	ppm	308.50	349.30	293.70	590.40	525.60	303.30	1826.00	1895.00	2070.00	934.30	3319.00	1214.00
Cs	ppm	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.
Cu	ppm	66.84	11.68	87.36	564.10	697.30	19.05	874.20	450.80	1540.00	504.20	1609.00	605.20
Dy	ppm	4.72	2.55	0.48	0.79	0.75	0.30	1.40	0.44	1.38	1.68	0.11	0.92
Er	ppm	2.89	1.65	0.26	0.47	0.45	0.21	0.80	0.27	0.81	1.05	0.07	0.59
Eu	ppm	1.09	0.48	0.12	0.14	0.13	0.04	0.23	0.18	0.19	0.10	0.02	0.05
Ga	ppm	13.96	11.95	3.46	5.99	5.77	3.93	8.09	4.23	8.36	11.42	5.84	7.08
Gd	ppm	3.93	1.88	0.43	0.63	0.60	0.25	1.21	0.38	1.12	1.32	0.09	0.75
Ge	ppm	1.85	0.75	0.50	0.43	0.40	0.28	1.57	1.64	1.95	0.61	5.09	0.74
Hf	ppm	2.45	0.97	0.27	0.48	0.45	0.28	0.75	0.22	0.68	0.75	< D.L.	0.39
Ho	ppm	1.05	0.59	0.10	0.18	0.16	0.07	0.31	0.10	0.30	0.39	0.02	0.22
ln La	ppm	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	0.11	< D.L.	0.18	< D.L.	< D.L.	< D.L.
La	ppm	5.40	0.99	0.80	0.83	0.83	0.44	2.26	0.61	1.55	1.33	< D.L.	0.55
Lu	ppm	0.43	0.25	0.04	0.07	0.07	0.04	0.12	0.04	0.12	0.15	0.01	0.09
Mo	ppm	< D.L.	< D.L.	5.58	12.11	8.58	1.43	< D.L.	< D.L.	< D.L.	1.39	184.30	14.06

Nb	ppm	6.84	0.57	1.08	1.46	1.47	0.89	3.90	0.88	2.96	2.07	< D.L.	1.16
Nd	ppm	10.30	3.43	1.28	1.60	1.52	0.89	3.45	1.01	2.69	2.59	0.21	1.20
Ni	ppm	112.80	126.40	140.40	492.70	385.90	190.70	1080.00	1180.00	1064.00	429.50	1609.00	710.70
Pb	ppm	0.90	< D.L.	< D.L.	1.23	1.17	1.08	< D.L.	< D.L.	< D.L.	< D.L.	2.24	1.22
Pr	ppm	2.05	0.58	0.26	0.31	0.29	0.18	0.75	0.21	0.56	0.49	0.04	0.20
Rb	ppm	0.79	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.	< D.L.					
Sc	ppm	38.58	32.40	5.40	8.67	8.67	5.12	15.97	9.30	16.01	22.31	9.47	12.85
Sb	ppm	< D.L.	< D.L.	< D.L.	< D.L.	0.25	< D.L.						
Sm	ppm	3.30	1.36	0.37	0.51	0.50	0.28	1.03	0.31	0.90	0.95	0.10	0.50
Sn	ppm	1.55	0.64	3.80	0.48	0.52	9.66	< D.L.	< D.L.	0.54	0.79	0.83	1.09
Sr	ppm	145.50	32.01	< D.L.	4.17	< D.L.	3.86	5.94	4.84	6.93	9.61	4.48	6.18
Та	ppm	0.52	0.05	0.09	0.11	0.11	0.07	0.28	0.06	0.21	0.16	< D.L.	0.08
Tb	ppm	0.69	0.35	0.07	0.11	0.11	0.04	0.21	0.06	0.20	0.24	0.02	0.14
Th	ppm	0.55	< D.L.	< D.L.	0.12	0.11	0.07	0.29	0.07	0.23	0.16	< D.L.	0.08
Tm	ppm	0.41	0.24	0.04	0.07	0.06	0.03	0.12	0.04	0.12	0.15	0.01	0.08
U	ppm	0.19	< D.L.	0.07	0.07	0.06	0.05	0.13	0.07	0.15	0.14	0.45	0.14
V	ppm	266.50	180.50	35.06	58.83	54.12	34.71	99.12	49.06	99.92	115.90	42.02	68.29
W	ppm	0.32	< D.L.	0.72	< D.L.	< D.L.	0.55	0.32	< D.L.	0.41	1.28	< D.L.	1.21
Υ	ppm	26.89	14.83	2.43	4.39	4.12	1.91	7.95	2.65	7.71	10.34	0.61	5.87
Yb	ppm	2.80	1.63	0.23	0.45	0.43	0.24	0.79	0.28	0.80	1.01	0.08	0.57
Zn	ppm	60.02	105.60	45.73	56.51	64.18	33.78	139.50	115.00	161.80	237.90	2947.00	379.70
Zr	ppm	84.25	29.36	10.77	17.07	16.19	10.14	27.35	7.70	24.15	24.61	< D.L.	14.00

<D.L.: lower than detection limit. Qz: quartz. Highly/moderately Si: highly/moderately silicified</p>

Table S2. Chemical analysis (wt.%) and structural formulas of chlorite, and calculated temperature of formation.

Sample	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195
Clast vs matrix	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Ma
Analysis #	5/1.	6/1.	9/1.	10 / 1	29 / 1	31 / 1	34 / 1	36 / 1	37 / 1	12 / 1	15 / 1	24 / 1	27 / 1	2/1.	3/1.	4/1.	5/1.	7/1.	8/1.	9/1.	10/1	11/1	12,
Chlorite composition a																							
SiO ₂	28.30	28.08	29.32	29.98	30.82	29.17	28.24	28.74	30.86	29.18	29.87	29.07	29.23	31.32	29.51	30.67	30.05	30.82	30.91	30.18	30.26	31.25	30.9
TiO ₂	0.01	0.04	0.02	0.00	0.01	0.01	0.06	0.00	0.02	0.01	0.00	0.00	0.08	0.02	0.00	0.00	0.04	0.00	0.04	0.00	0.01	0.02	0.0
Al ₂ O ₃	19.56	19.77	18.45	18.28	17.94	18.94	20.21	19.39	17.39	19.45	19.06	19.14	18.93	18.64	18.77	17.80	17.65	18.54	17.51	18.48	17.67	17.83	18.
Cr ₂ O ₃	0.00	0.00	0.00	0.00	0.01	0.00	0.06	0.05	0.00	0.00	0.00	0.07	0.00	0.00	0.00	0.01	0.01	0.03	0.08	0.03	0.04	0.04	0.0
FeO	18.76	18.44	16.25	15.57	16.08	16.45	15.93	15.62	14.80	16.77	16.88	16.86	16.09	14.84	14.91	15.42	16.29	14.63	14.77	15.25	14.64	15.73	14.
MnO	0.49	0.37	0.32	0.25	0.39	0.38	0.19	0.23	0.31	0.43	0.38	0.35	0.36	0.35	0.42	0.37	0.44	0.34	0.39	0.33	0.46	0.37	0.3
MgO	19.38	19.96	22.28	22.99	22.03	21.12	22.20	22.85	23.26	21.70	21.47	21.86	21.58	23.24	22.82	21.58	21.48	23.53	22.65	23.16	23.04	22.40	23.
NiO	0.07	0.00	0.02	0.04	0.02	0.07	0.01	0.00	0.04	0.02	0.12	0.01	0.05	0.00	0.06	0.05	0.02	0.07	0.03	0.02	0.00	0.05	0.0
CaO	0.05	0.07	0.19	0.10	0.14	0.14	0.09	0.05	0.11	0.14	0.15	0.14	0.20	0.21	0.16	0.25	0.21	0.16	0.25	0.16	0.20	0.23	0.2
Na₂O	0.05	0.02	0.11	0.06	0.05	0.05	0.02	0.03	0.05	0.02	0.03	0.04	0.05	0.06	0.06	0.03	0.07	0.08	0.08	0.05	0.10	0.09	0.0
K ₂ O	0.00	0.02	0.05	0.06	0.05	0.04	0.03	0.02	0.03	0.01	0.03	0.02	0.04	0.06	0.05	0.06	0.13	0.08	0.10	0.05	0.06	0.09	0.0
total	86.68	86.77	87.00	87.32	87.54	86.37	87.04	86.97	86.87	87.74	87.99	87.54	86.62	88.73	86.75	86.26	86.40	88.30	86.80	87.72	86.48	88.09	87.
K ₂ O+Na ₂ O+CaO	0.10	0.11	0.35	0.21	0.24	0.23	0.14	0.10	0.19	0.17	0.21	0.20	0.29	0.32	0.27	0.34	0.42	0.32	0.43	0.26	0.36	0.41	0.3
Structural formula b																							
Si	2.91	2.87	2.96	3.00	3.08	2.97	2.84	2.89	3.09	2.92	2.98	2.92	2.96	3.06	2.96	3.10	3.05	3.03	3.10	3.00	3.04	3.09	3.0
Ti	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.0
Al	2.37	2.38	2.19	2.15	2.11	2.27	2.40	2.30	2.05	2.30	2.24	2.27	2.26	2.15	2.22	2.12	2.11	2.15	2.07	2.16	2.10	2.08	2.
Al ^{IV}	1.09	1.13	1.04	1.00	0.92	1.03	1.16	1.11	0.91	1.08	1.02	1.08	1.04	0.94	1.04	0.90	0.95	0.97	0.90	1.00	0.96	0.91	0.9
Al ^{vi}	1.27	1.26	1.15	1.15	1.19	1.24	1.24	1.19	1.13	1.22	1.23	1.19	1.22	1.21	1.19	1.22	1.17	1.18	1.16	1.16	1.14	1.18	1.3
Cr	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.0
Fe	1.61	1.58	1.37	1.30	1.34	1.40	1.34	1.31	1.24	1.41	1.41	1.42	1.36	1.21	1.25	1.30	1.38	1.20	1.24	1.27	1.23	1.30	1.2
Mn	0.04	0.03	0.03	0.02	0.03	0.03	0.02	0.02	0.03	0.04	0.03	0.03	0.03	0.03	0.04	0.03	0.04	0.03	0.03	0.03	0.04	0.03	0.0
Mg	2.97	3.05	3.35	3.43	3.28	3.20	3.33	3.43	3.47	3.24	3.20	3.28	3.26	3.39	3.42	3.25	3.25	3.45	3.38	3.43	3.46	3.31	3.4
Ni	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.0
Са	0.01	0.01	0.02	0.01	0.01	0.02	0.01	0.00	0.01	0.01	0.02	0.01	0.02	0.02	0.02	0.03	0.02	0.02	0.03	0.02	0.02	0.02	0.0
Na	0.01	0.00	0.02	0.01	0.01	0.01	0.00	0.01	0.01	0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.02	0.01	0.01	0.02	0.02	0.0
K	0.00	0.00	0.01	0.01	0.01	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.01	0.02	0.01	0.01	0.01	0.01	0.01	0.0
octaedral sum	5.91	5.93	5.95	5.93	5.87	5.91	5.95	5.96	5.89	5.93	5.90	5.95	5.91	5.87	5.93	5.85	5.90	5.91	5.88	5.93	5.92	5.87	5.8
vacancies	0.09	0.07	0.05	0.07	0.13	0.09	0.05	0.04	0.11	0.07	0.10	0.05	0.09	0.13	0.07	0.15	0.10	0.09	0.12	0.07	0.08	0.13	0.
Mg# ^c	0.65	0.66	0.71	0.72	0.71	0.70	0.71	0.72	0.74	0.70	0.69	0.70	0.71	0.74	0.73	0.71	0.70	0.74	0.73	0.73	0.74	0.72	0.7
R ^{2+ d}	4.63	4.66	4.75	4.75	4.66	4.64	4.69	4.76	4.73	4.69	4.65	4.72	4.65	4.63	4.71	4.59	4.68	4.69	4.65	4.73	4.73	4.64	4.
Calculated T (°C)																							
Bourdelle et al. (2013) ^e Tcorrected (>350°C) ^e	261	300	318	269	197	238	373 <i>324</i>	393 <i>332</i>	210	282	230	321	248	198	273	178	217	228	196	260	241	192	20

^a EPMA analysis of chlorite composition in wt.% of oxides, selected if oxide sum is in the 86-89 wt.% range and Na₂O + K₂O + CaO < 1 wt.%

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^b Chlorite structural formula calculated on a 14-oxygen basis. Selection criterias: Si ≤ 4, cation sum ≤ 10.1 and vacancies > 0.01

 $^{^{}c}$ Mg# = Mg / (Mg+Fe)

 $^{^{}d} R^{2+} = (Fe + Mg + Mn + Ni)$

^e temperature of chlorite formation calculated from R²⁺ and Si with the thermometer of Bourdelle et al. (2013) calibrated for Tchl < 350 °C; a quadratic correction is applied for Tchl > 350 °C (Tcorrected)

Sample	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195	195
Clast vs matrix	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat
Analysis #	13 / 1	17 / 1	18 / 1	19 / 1	20 / 1	21/1	22 / 1	23 / 1	24 / 1	25 / 1	26 / 1	28 / 1	30/1	31/1	32 / 1	33 / 1	34 / 1	35 / 1	36 / 1	37 / 1	38 / 1	39 / 1	40 / 1
Chlorite composition a																							
SiO ₂	30.06	30.55	30.97	30.15	31.84	30.68	30.56	31.26	30.86	30.25	30.44	30.43	31.32	30.78	29.48	29.68	29.70	30.14	28.72	29.06	29.12	29.76	29.65
TiO ₂	0.02	0.01	0.01	0.02	0.03	0.04	0.00	0.04	0.00	0.02	0.08	0.00	0.00	0.00	0.00	0.00	0.02	0.00	0.07	0.04	0.00	0.02	0.07
Al_2O_3	18.67	18.30	17.70	17.85	18.36	17.95	18.30	17.65	18.58	18.66	18.48	17.69	17.71	18.17	18.76	18.91	18.65	18.16	19.05	19.18	18.94	18.71	18.58
Cr ₂ O ₃	0.04	0.01	0.04	0.07	0.01	0.02	0.02	0.00	0.00	0.08	0.01	0.05	0.00	0.01	0.04	0.01	0.01	0.00	0.00	0.04	0.03	0.00	0.00
FeO	14.62	13.98	15.61	14.03	15.44	14.95	14.36	15.08	13.67	14.59	14.52	14.41	14.23	13.18	17.42	16.90	17.56	16.12	17.77	17.01	18.17	17.79	15.36
MnO	0.45	0.29	0.44	0.23	0.45	0.38	0.29	0.41	0.42	0.23	0.36	0.31	0.38	0.31	0.30	0.37	0.40	0.40	0.34	0.43	0.40	0.43	0.31
MgO	22.47	24.25	21.86	23.75	21.73	22.86	23.18	22.53	24.23	24.82	22.99	24.49	23.73	25.12	21.68	21.72	22.16	22.11	21.80	21.58	20.68	20.87	22.73
NiO	0.03	0.12	0.06	0.08	0.03	0.00	0.05	0.00	0.01	0.13	0.00	0.08	0.10	0.01	0.04	0.03	0.01	0.00	0.01	0.04	0.01	0.06	0.01
CaO	0.24	0.19	0.17	0.13	0.33	0.26	0.27	0.30	0.06	0.10	0.15	0.13	0.17	0.11	0.14	0.13	0.12	0.21	0.08	0.10	0.15	0.17	0.14
Na₂O	0.02	0.03	0.05	0.03	0.03	0.04	0.06	0.04	0.06	0.05	0.05	0.02	0.06	0.06	0.04	0.02	0.06	0.07	0.00	0.02	0.02	0.01	0.05
K ₂ O	0.05	0.01	0.06	0.03	0.09	0.04	0.04	0.06	0.05	0.02	0.02	0.05	0.06	0.05	0.02	0.04	0.03	0.07	0.03	0.02	0.04	0.06	0.03
total	86.68	87.73	86.96	86.39	88.34	87.22	87.13	87.36	87.95	88.96	87.09	87.67	87.75	87.79	87.93	87.81	88.71	87.29	87.86	87.51	87.54	87.87	86.95
K ₂ O+Na ₂ O+CaO	0.31	0.23	0.28	0.20	0.46	0.34	0.37	0.39	0.17	0.17	0.21	0.20	0.28	0.21	0.20	0.19	0.20	0.36	0.11	0.13	0.20	0.24	0.22
Structural formula b																							
Si	3.01	3.01	3.11	3.02	3.13	3.06	3.04	3.11	3.03	2.96	3.03	3.02	3.09	3.02	2.96	2.97	2.96	3.02	2.89	2.93	2.95	2.99	2.98
Ti	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.01
Al	2.21	2.13	2.09	2.11	2.13	2.11	2.15	2.07	2.15	2.15	2.17	2.07	2.06	2.10	2.22	2.23	2.19	2.15	2.26	2.28	2.26	2.22	2.20
Al ^{IV}	0.99	0.99	0.89	0.98	0.87	0.94	0.96	0.89	0.97	1.04	0.97	0.98	0.91	0.98	1.04	1.03	1.04	0.98	1.11	1.07	1.05	1.01	1.02
Al ^{VI}	1.22	1.14	1.20	1.13	1.26	1.17	1.19	1.18	1.18	1.10	1.20	1.08	1.15	1.12	1.18	1.20	1.14	1.17	1.15	1.20	1.21	1.21	1.17
Cr	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Fe	1.23	1.15	1.31	1.18	1.27	1.25	1.20	1.26	1.12	1.19	1.21	1.19	1.17	1.08	1.46	1.41	1.46	1.35	1.50	1.43	1.54	1.50	1.29
Mn	0.04	0.02	0.04	0.02	0.04	0.03	0.02	0.03	0.03	0.02	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.04	0.03	0.04	0.03
Mg	3.36	3.57	3.27	3.55	3.19	3.40	3.44	3.34	3.54	3.62	3.41	3.62	3.49	3.68	3.24	3.24	3.29	3.31	3.27	3.24	3.12	3.13	3.40
Ni	0.00	0.01	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.01	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00
Ca	0.03	0.02	0.02	0.01	0.04	0.03	0.03	0.03	0.01	0.01	0.02	0.01	0.02	0.01	0.01	0.01	0.01	0.02	0.01	0.01	0.02	0.02	0.02
Na	0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.00	0.01	0.01	0.01	0.00	0.01	0.01	0.00	0.00	0.00	0.00	0.01
K	0.01	0.00	0.01	0.00	0.01	0.00	0.00	0.01	0.01	0.00	0.00	0.01	0.01	0.01	0.00	0.01	0.00	0.01	0.00	0.00	0.00	0.01	0.00
octaedral sum	5.88	5.92	5.85	5.92	5.81	5.89	5.89	5.86	5.91	5.97	5.88	5.95	5.89	5.94	5.94	5.92	5.95	5.91	5.97	5.93	5.93	5.90	5.92
vacancies	0.12	0.08	0.15	0.08	0.19	0.11	0.11	0.14	0.09	0.03	0.12	0.05	0.11	0.06	0.06	0.08	0.05	0.09	0.03	0.07	0.07	0.10	0.08
Mg# c	0.73	0.76	0.71	0.75	0.71	0.73	0.74	0.73	0.76	0.75	0.74	0.75	0.75	0.77	0.69	0.70	0.69	0.71	0.69	0.69	0.67	0.68	0.73
R ^{2+ d}	4.62	4.75	4.62	4.75	4.49	4.68	4.66	4.63	4.70	4.84	4.65	4.84	4.70	4.78	4.73	4.69	4.78	4.70	4.80	4.71	4.69	4.67	4.72
Calculated T (°C)													3	2			3	3		=		****	
Bourdelle et al. (2013) e																							
Tcorrected (>350°C) e	216	255	180	252	156	211	216	183	228	403	211	319	203	272	290	254	333	236	443	294	274	233	265
, ,																							

Sample	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155
Clast vs matrix	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast
Analysis #	14	73 / 1	75 / 1	76 / 1	77 / 1	81 / 1	82 / 1	83 / 1	41 / 1	42 / 1	43 / 1	44 / 1	45 / 1	46 / 1	48 / 1	49 / 1	50 / 1	51 / 1	52 / 1	53 / 1	54 / 1	57 / 1	58 / 1
Chlorite composition ^a																							
SiO ₂	28.07	31.03	30.61	30.78	30.68	29.35	27.02	29.24	27.94	28.79	26.59	26.61	25.39	28.07	29.40	29.06	28.72	27.25	29.02	28.86	29.17	27.82	26.49
TiO₂	0.03	0.04	0.08	0.06	0.03	0.01	0.07	0.02	0.00	0.02	0.05	0.06	0.02	0.01	0.00	0.04	0.09	0.07	0.00	0.01	0.01	0.05	0.05
Al ₂ O ₃	19.16	19.99	18.27	20.81	18.41	17.26	19.23	17.37	18.57	18.11	19.93	19.88	18.88	18.00	18.17	18.03	17.80	19.41	17.68	18.03	17.46	20.26	20.36
Cr ₂ O₃	0.03	0.08	0.01	0.00	0.06	0.00	0.10	0.02	0.11	0.21	0.08	0.04	0.00	0.08	0.26	0.40	0.44	0.04	0.31	0.43	0.30	0.12	0.00
FeO	22.98	19.31	19.62	19.95	20.75	22.31	24.31	21.62	22.64	24.05	25.12	25.55	29.02	21.98	21.13	21.07	22.37	24.60	22.35	22.81	22.68	24.06	25.08
MnO	0.27	0.35	0.31	0.53	0.33	0.32	0.39	0.24	0.23	0.41	0.24	0.29	0.17	0.52	0.56	0.52	0.38	0.39	0.33	0.35	0.21	0.21	0.32
MgO	17.21	15.68	17.35	14.91	15.33	18.53	16.82	18.95	17.08	16.84	15.17	15.82	12.40	18.19	17.25	16.60	17.12	15.04	16.17	16.74	16.60	15.22	14.27
NiO	0.09	0.09	0.07	0.15	0.12	0.11	0.00	0.09	0.14	0.08	0.00	0.01	0.03	0.09	0.01	0.17	0.08	0.05	0.07	0.17	0.06	0.00	0.02
CaO	0.09	0.18	0.22	0.23	0.30	0.22	0.06	0.09	0.07	0.10	0.07	0.07	0.07	0.10	0.11	0.11	0.09	0.05	0.27	0.13	0.18	0.08	0.11
Na₂O	0.01	0.08	0.06	0.08	0.11	0.01	0.06	0.04	0.00	0.03	0.04	0.07	0.06	0.04	0.07	0.10	0.05	0.06	0.12	0.07	0.04	0.05	0.03
K₂O	0.01	0.02	0.04	0.05	0.02	0.00	0.02	0.00	0.01	0.00	0.01	0.02	0.01	0.02	0.05	0.05	0.01	0.02	0.04	0.01	0.01	0.02	0.02
total	87.94	86.85	86.63	87.56	86.15	88.12	88.07	87.68	86.79	88.65	87.30	88.41	86.06	87.09	87.00	86.15	87.16	86.99	86.35	87.60	86.72	87.89	86.74
K ₂ O+Na ₂ O+CaO	0.10	0.28	0.32	0.36	0.43	0.23	0.14	0.12	0.08	0.14	0.12	0.15	0.14	0.17	0.22	0.26	0.15	0.13	0.43	0.21	0.23	0.15	0.16
Structural formula b																							
Si	2.90	3.15	3.14	3.12	3.18	3.02	2.82	3.01	2.93	2.97	2.81	2.78	2.79	2.93	3.04	3.04	2.99	2.88	3.05	3.00	3.05	2.89	2.82
Ti	0.00	0.00	0.01	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.00	0.00	0.00
Al	2.34	2.40	2.21	2.48	2.25	2.09	2.37	2.11	2.29	2.20	2.48	2.45	2.45	2.21	2.22	2.22	2.19	2.42	2.19	2.21	2.15	2.48	2.55
Al ^{IV}	1.10	0.85	0.86	0.88	0.82	0.98	1.18	0.99	1.07	1.03	1.19	1.22	1.21	1.07	0.96	0.96	1.01	1.12	0.95	1.00	0.95	1.11	1.18
Al ^{vi}	1.24	1.55	1.35	1.60	1.43	1.11	1.19	1.12	1.22	1.17	1.29	1.23	1.24	1.14	1.26	1.27	1.18	1.30	1.24	1.21	1.21	1.37	1.37
Cr	0.00	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.01	0.02	0.01	0.00	0.00	0.01	0.02	0.03	0.04	0.00	0.03	0.04	0.02	0.01	0.00
Fe	1.99	1.64	1.68	1.69	1.80	1.92	2.12	1.86	1.98	2.08	2.22	2.24	2.67	1.92	1.83	1.84	1.95	2.17	1.96	1.98	1.99	2.09	2.23
Mn	0.02	0.03	0.03	0.05	0.03	0.03	0.03	0.02	0.02	0.04	0.02	0.03	0.02	0.05	0.05	0.05	0.03	0.04	0.03	0.03	0.02	0.02	0.03
Mg	2.65	2.38	2.65	2.25	2.37	2.84	2.62	2.91	2.67	2.59	2.39	2.47	2.03	2.83	2.66	2.59	2.66	2.37	2.53	2.59	2.59	2.36	2.26
Ni	0.01	0.01	0.01	0.01	0.01	0.01	0.00	0.01	0.01	0.01	0.00	0.00	0.00	0.01	0.00	0.01	0.01	0.00	0.01	0.01	0.00	0.00	0.00
Са	0.01	0.02	0.02	0.03	0.03	0.02	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.03	0.01	0.02	0.01	0.01
Na	0.00	0.02	0.01	0.01	0.02	0.00	0.01	0.01	0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.02	0.01	0.01	0.02	0.01	0.01	0.01	0.01
K	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.01	0.00	0.00	0.00	0.00
octaedral sum	5.92	5.65	5.75	5.64	5.70	5.93	5.99	5.94	5.92	5.92	5.94	5.99	5.99	5.97	5.85	5.84	5.89	5.91	5.86	5.89	5.86	5.86	5.91
vacancies	0.08	0.35	0.25	0.36	0.30	0.07	0.01	0.06	0.08	0.08	0.06	0.01	0.01	0.03	0.15	0.16	0.11	0.09	0.14	0.11	0.14	0.14	0.09
Mg# ^c	0.57	0.59	0.61	0.57	0.57	0.60	0.55	0.61	0.57	0.56	0.52	0.52	0.43	0.60	0.59	0.58	0.58	0.52	0.56	0.57	0.57	0.53	0.50
R ^{2+ d}	4.67	4.06	4.37	4.00	4.21	4.80	4.77	4.80	4.68	4.71	4.63	4.73	4.72	4.80	4.54	4.50	4.65	4.58	4.53	4.62	4.60	4.46	4.52
Calculated T (°C)																							
Bourdelle et al. (2013) e	284	113	138	116	120	274	1305	278	276	272	359	863	612	399	192	185	243	262	193	231	200	223	273
Tcorrected (>350°C) e							492				317	444	398	334									

Sample	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a	218a
Clast vs matrix	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast
Analysis #	22	29	30	13 / 1	14 / 1	18 / 1	19 / 1	20 / 1	21 / 1	23 / 1	26 / 1	27 / 1	33 / 1	39 / 1	41/1	43 / 1	49 / 1	56/1	57 / 1	58 / 1	59 / 1	60 / 1	61 / 1
Zone analysed	C2	C6	C6	C-3	C-4	C-4	C-4	C-4	C-4	C-4	C-5	C-6	C-6	C-6	C-7	C-7	C-7						
Chlorite composition a																							
SiO ₂	27.59	30.42	27.21	29.33	28.53	27.11	27.18	28.94	25.64	26.69	29.01	28.37	29.10	28.70	27.66	28.07	28.88	29.91	30.98	29.13	28.05	27.65	28.23
TiO ₂	0.03	0.02	0.03	0.00	0.08	0.06	0.11	0.06	0.04	0.06	0.00	0.08	0.00	0.34	0.04	0.11	0.03	0.03	0.04	0.01	0.07	0.06	0.09
Al ₂ O ₃	19.06	19.76	19.12	17.56	17.23	19.43	18.92	18.86	19.64	19.08	19.64	19.59	17.66	18.09	18.48	18.21	17.55	17.48	15.91	18.13	19.30	19.37	19.31
Cr ₂ O ₃	0.03	0.04	0.05	0.07	0.07	0.08	0.19	0.00	0.04	0.12	0.04	0.08	0.17	0.05	0.17	0.12	0.16	0.66	0.64	0.35	0.09	0.05	0.01
FeO	23.76	9.81	24.59	23.27	23.60	25.89	23.53	18.33	28.21	26.53	18.58	17.75	22.42	20.09	21.23	21.92	18.37	17.41	14.20	14.87	18.70	19.11	18.76
MnO	0.42	0.25	0.47	0.58	0.45	0.51	0.59	0.73	0.37	0.50	0.21	0.18	0.44	0.57	0.35	0.54	0.30	0.01	0.06	0.11	0.23	0.07	0.20
MgO	17.14	27.78	16.45	16.29	16.29	13.86	15.44	21.24	12.65	13.98	20.59	21.11	17.87	17.73	18.06	16.77	21.61	21.13	25.20	23.31	19.73	20.50	20.66
NiO	0.01	0.10	0.00	0.00	0.11	0.00	0.02	0.04	0.09	0.00	0.00	0.08	0.05	0.06	0.06	0.11	0.08	0.23	0.26	0.23	0.02	0.05	0.00
CaO	0.13	0.03	0.09	0.23	0.24	0.15	0.26	0.07	0.05	0.11	0.04	0.11	0.17	0.46	0.15	0.23	0.09	0.07	0.01	0.02	0.19	0.03	0.03
Na₂O	0.04	0.02	0.03	0.00	0.02	0.04	0.01	0.00	0.05	0.02	0.01	0.00	0.02	0.01	0.03	0.00	0.03	0.00	0.00	0.00	0.06	0.03	0.01
K ₂ O	0.01	0.02	0.01	0.03	0.02	0.01	0.01	0.02	0.03	0.00	0.00	0.00	0.01	0.01	0.00	0.04	0.01	0.00	0.00	0.04	0.05	0.02	0.01
total	88.23	88.23	88.06	87.37	86.64	87.14	86.26	88.29	86.82	87.10	88.10	87.34	87.92	86.10	86.24	86.13	87.11	86.94	87.33	86.21	86.48	86.93	87.31
K₂O+Na₂O+CaO	0.18	0.07	0.14	0.26	0.28	0.20	0.28	0.09	0.13	0.13	0.05	0.11	0.20	0.48	0.18	0.27	0.13	0.08	0.02	0.06	0.29	0.08	0.05
Structural formula ^b																							
Si	2.86	2.92	2.84	3.06	3.01	2.88	2.89	2.91	2.78	2.85	2.92	2.87	3.00	2.99	2.90	2.96	2.95	3.04	3.08	2.95	2.89	2.84	2.88
Ti	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.03	0.00	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.01
Al	2.33	2.24	2.36	2.16	2.15	2.43	2.37	2.24	2.51	2.40	2.33	2.34	2.15	2.22	2.28	2.26	2.11	2.09	1.87	2.16	2.34	2.34	2.32
Aliv	1.14	1.08	1.16	0.94	0.99	1.12	1.11	1.09	1.22	1.15	1.08	1.13	1.00	1.01	1.10	1.04	1.05	0.96	0.92	1.05	1.11	1.16	1.12
Al ^{vi}	1.19	1.16	1.20	1.21	1.16	1.32	1.26	1.15	1.29	1.26	1.24	1.21	1.15	1.22	1.19	1.22	1.06	1.13	0.95	1.11	1.23	1.18	1.19
Cr	0.00	0.00	0.00	0.01	0.01	0.01	0.02	0.00	0.00	0.01	0.00	0.01	0.01	0.00	0.01	0.01	0.01	0.05	0.05	0.03	0.01	0.00	0.00
Fe	2.06	0.79	2.15	2.03	2.09	2.30	2.09	1.54	2.56	2.37	1.56	1.50	1.94	1.75	1.86	1.93	1.57	1.48	1.18	1.26	1.61	1.64	1.60
Mn	0.04	0.02	0.04	0.05	0.04	0.05	0.05	0.06	0.03	0.04	0.02	0.02	0.04	0.05	0.03	0.05	0.03	0.00	0.01	0.01	0.02	0.01	0.02
Mg	2.65	3.98	2.56	2.53	2.57	2.20	2.45	3.19	2.05	2.23	3.09	3.19	2.75	2.76	2.82	2.64	3.29	3.20	3.74	3.52	3.03	3.14	3.14
Ni	0.00	0.01	0.00	0.00	0.01	0.00	0.00	0.00	0.01	0.00	0.00	0.01	0.00	0.01	0.01	0.01	0.01	0.02	0.02	0.02	0.00	0.00	0.00
Ca	0.01	0.00	0.01	0.03	0.03	0.02	0.03	0.01	0.01	0.01	0.00	0.01	0.02	0.05	0.02	0.03	0.01	0.01	0.00	0.00	0.02	0.00	0.00
Na	0.01	0.00	0.01	0.00	0.00	0.01	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.01	0.00	0.00	0.00	0.01	0.01	0.00
K	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.01	0.01	0.00	0.00
octaedral sum	5.97	5.96	5.98	5.86	5.90	5.89	5.90	5.96	5.96	5.93	5.92	5.94	5.92	5.84	5.95	5.89	5.99	5.89	5.95	5.96	5.94	5.98	5.95
vacancies	0.03	0.04	0.02	0.14	0.10	0.11	0.10	0.04	0.04	0.07	0.08	0.06	0.08	0.16	0.05	0.11	0.01	0.11	0.05	0.04	0.06	0.02	0.05
Mg# c	0.56	0.83	0.54	0.56	0.55	0.49	0.54	0.67	0.44	0.48	0.66	0.68	0.59	0.61	0.60	0.58	0.68	0.68	0.76	0.74	0.65	0.66	0.66
R ^{2+ d}	4.75	4.79	4.75	4.61	4.70	4.54	4.59	4.80	4.65	4.65	4.67	4.71	4.73	4.57	4.72	4.63	4.89	4.70	4.95	4.81	4.66	4.79	4.75
Calculated T (°C)																							
Bourdelle et al. (2013) e	445	362	500	194	234	247	263	388	407	313	271	336	251	203	330	233	585	211	302	330	304	630	379
Tcorrected (>350°C) ^e	351	319	369					330	337								391					402	326

Sample Clast vs matrix Analysis # 218a clast matrix Analysis # 218a clast clast clast clast clast clast matrix clast clast matrix matrix follows and matrix matrix follows and matrix matrix follows and mat	
Analysis # 62 /1 63 /1 64 /1 65 /1 9 /1 10 /1 11 /1 44 /1 45 /1 46 /1 47 /1 50 /1 Chlorite composition ** SiO₂ 28.23 29.35 27.75 28.58 28.20 30.04 28.31 27.60 28.93 28.93 30.16 TiO₂ 0.01 0.08 0.04 0.04 0.00 0.03 0.00 0.04 0.00 0.09 0.03 0.00 0.00 0.09 0.03 0.00 Al₂O₃ 19.86 18.85 20.29 20.27 18.42 17.90 18.64 18.47 19.06 19.56 18.66 Cr₂O₃ 10.00 0.01 0.02 0.01 0.12 0.22 20.93 21.55 21.50 18.64 18.47 19.06 19.56 18.66 MiO 0.16 0.07 0.18 0.08 0.40 0.16 0.40 0.15 0.21 0.02 0.02 0.03 0.00 0.01	218a
Chlorite composition ^a SiO ₂ 28.23 29.35 27.75 28.58 28.20 30.04 28.31 27.60 28.93 28.93 28.65 30.16 TiO₂ 0.01 0.08 0.04 0.04 0.00 0.03 0.00 0.03 0.00 0.09 0.03 0.00 Al₂O₃ 19.86 18.85 20.29 20.27 18.42 17.92 17.90 18.64 18.47 19.06 19.56 18.66 Cr₂O₃ 0.00 0.01 0.02 0.11 0.27 0.67 0.08 0.11 0.03 0.04 0.26 0.17 FeO 17.36 17.43 18.52 18.22 29.93 22.55 21.15 23.60 19.10 14.22 18.97 17.54 MnO 0.16 0.17 0.18 0.80 0.40 0.61 0.25 0.18 0.40 0.25 0.28 0.25 0.20 0.28 0.25 0.28 0.00 0.01 <t< th=""><th>Mat</th></t<>	Mat
SiO₂ 28.23 29.35 27.75 28.58 28.20 30.04 28.31 27.60 28.93 28.98 28.65 30.16 TiO₂ 0.01 0.08 0.04 0.04 0.00 0.03 0.00 0.04 0.00 0.03 0.00 0.04 0.00 0.03 0.00 0.04 0.00 0.03 0.00 0.04 0.00 0.03 0.00 Al₂O₃ 19.86 18.85 20.29 20.27 18.42 17.92 17.90 18.64 18.47 19.06 19.56 18.66 0.17 0.67 0.08 0.11 0.03 0.04 0.26 0.17 75.4 MnO 0.16 0.17 0.15 0.21 0.38 0.40 0.36 0.43 0.25 0.28 0.26 0.25 MgO 2.17 2.100 2.083 19.95 18.08 15.79 18.24 16.47 20.11 20.25 0.24 0.24 0.04 0.15 <th< th=""><th>69 / 1</th></th<>	69 / 1
TiO₂ 0.01 0.08 0.04 0.04 0.00 0.03 0.00 0.04 0.00 0.09 0.03 0.00 Al₂O₃ 19.86 18.85 20.29 20.27 18.42 17.90 18.64 18.47 19.06 19.56 18.66 C₂O₃ 0.00 0.01 0.02 0.11 0.27 0.08 0.11 0.03 0.04 0.26 0.17 FeO 17.36 17.43 18.52 18.22 22.93 22.55 21.15 23.60 19.10 14.22 18.97 17.54 MgO 21.71 21.00 20.83 19.95 18.08 15.79 18.24 16.47 20.11 23.98 19.42 20.42 MiO 0.02 0.07 0.16 0.06 0.05 0.08 0.10 0.05 0.08 0.03 0.00 0.01 0.02 A₂O 0.00 0.01 0.01 0.00 0.00 0.00 0.00 0.00<	
Al₂O₃ 19.86 18.85 20.29 20.27 18.42 17.92 17.90 18.64 18.47 19.06 19.56 18.66 Cr₂O₃ 0.00 0.01 0.02 0.11 0.27 0.67 0.08 0.11 0.03 0.04 0.26 0.17 FeO 17.36 17.34 18.52 18.22 12.23 22.55 21.15 23.60 19.10 14.22 18.97 17.54 MnO 0.16 0.17 21.00 20.83 19.95 18.08 15.79 18.24 16.47 20.11 23.98 19.42 20.42 NiO 0.02 0.07 0.16 0.06 0.0 0.04 0.16 0.04 0.25 0.00 0.07 0.11 0.00 Argo 0.00 0.01 0.01 0.00 0.02 0.00 0.08 0.03 0.00 0.03 0.03 K2O+Na ₂ O+CaO 0.00 0.01 0.05 0.09 0.05	29.44
Cr₂O₃ 0.00 0.01 0.02 0.11 0.27 0.67 0.08 0.11 0.03 0.04 0.26 0.17 FeO 17.36 17.43 18.52 18.22 22.93 22.55 21.15 23.60 19.10 14.22 18.97 17.54 MnO 0.16 0.17 0.15 0.21 0.38 0.40 0.36 0.43 0.25 0.28 0.26 0.26 MgO 21.71 21.00 20.83 19.95 18.08 15.79 18.24 16.04 20.11 23.98 19.42 20.42 NiO 0.02 0.07 0.16 0.06 0.05 0.28 0.10 0.15 0.07 0.11 0.02 CaO 0.00 0.01 0.00 0.00 0.05 0.88 0.00 0.00 0.07 0.01 0.00 Na2O 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	0.03
FeO 17.36 17.43 18.52 18.22 22.93 22.55 21.15 23.60 19.10 14.22 18.97 17.54 MnO 0.16 0.17 0.15 0.21 0.38 0.40 0.36 0.43 0.25 0.28 0.26 0.26 MgO 21.71 21.00 20.83 19.95 18.08 15.79 18.24 16.47 20.11 23.98 19.42 20.42 NiO 0.02 0.07 0.16 0.06 0.10 0.04 0.15 0.04 0.25 0.00 0.01 0.02 CaO 0.00 0.01 0.00 0.05 0.28 0.08 0.03 0.00 0.02 0.05 K₂O 0.00 0.00 0.02 0.02 0.00 0.08 0.00 0.00 0.00 0.03 0.03 0.00 0.03 0.03 K₂O+Na₂O+CaO 0.00 0.01 0.00 0.05 0.49 0.18 0.18	18.49
MnO 0.16 0.17 0.15 0.21 0.38 0.40 0.36 0.43 0.25 0.28 0.26 0.26 MgO 21.71 21.00 20.83 19.95 18.08 15.79 18.24 16.47 20.11 23.98 19.42 20.42 NiO 0.02 0.07 0.16 0.06 0.10 0.04 0.16 0.04 0.25 0.10 0.11 0.02 CaO 0.00 0.01 0.01 0.00 0.00 0.05 0.28 0.10 0.15 0.00 0.07 0.11 0.05 K₂O 0.00 0.00 0.02 0.02 0.00 0.00 0.00 0.03 0.03 0.03 Structural formula b V <t< th=""><th>0.03</th></t<>	0.03
MgO 21.71 21.00 20.83 19.95 18.08 15.79 18.24 16.47 20.11 23.98 19.42 20.42 NiO 0.02 0.07 0.16 0.06 0.10 0.04 0.16 0.04 0.25 0.10 0.11 0.02 CaO 0.00 0.01 0.01 0.00 0.05 0.28 0.10 0.15 0.00 0.01 0.01 0.06 Na ₂ O 0.00	14.29
NiO 0.02 0.07 0.16 0.06 0.10 0.04 0.16 0.04 0.25 0.10 0.11 0.02 CaO 0.00 0.11 0.03 0.06 0.05 0.28 0.10 0.15 0.00 0.07 0.11 0.06 Na₂O 0.00 0.01 0.01 0.00 0.00 0.05 0.08 0.03 0.00 0.02 0.03 0.03 total 87.35 87.08 87.82 87.52 88.42 87.85 86.38 87.11 87.13 86.87 87.50 87.37 K₂O+Na₂O+CaO 0.00 0.11 0.05 0.09 0.05 0.40 0.18 0.18 0.00 0.12 0.23 0.13 Structural formula b 87.35 87.08 2.81 2.89 2.91 3.10 2.96 2.90 2.90 2.92 3.04 Ti 0.00 0.01 0.00 0.00 0.00 0.00 0.00 0.	0.11
CaO 0.00 0.11 0.03 0.06 0.05 0.28 0.10 0.15 0.00 0.07 0.11 0.06 Na₂O 0.00 0.01 0.00 0.02 0.02 0.00 0.08 0.03 0.00 0.02 0.03 0.03 total 87.35 87.08 87.82 87.52 88.42 87.85 86.38 87.11 87.13 86.87 87.50 87.37 K₂O+Na₂O+CaO 0.00 0.11 0.05 0.09 0.05 0.40 0.18 0.18 0.00 0.12 0.23 0.13 Structural formula b V V V V V V V V 0.00 0.01 0.00 <	23.76
Na₂O 0.00 0.01 0.01 0.00 0.00 0.05 0.08 0.03 0.00 0.02 0.03 k₂O 0.00 0.00 0.02 0.02 0.00 0.08 0.00 0.00 0.03 0.03 0.03 k₂O+Na₂O+CaO 0.00 0.11 0.05 0.09 0.05 0.40 0.18 0.18 0.00 0.12 0.23 0.13 Structural formula b V V V V V V V V V 0.00 0.18 0.18 0.00 0.12 0.23 0.13 Structural formula b V V V V V V V V V V V 0.00 0.10 0.00 <td>0.28</td>	0.28
K₂O 0.00 0.00 0.02 0.02 0.00 0.08 0.00 0.00 0.03 0.00 <th< th=""><th>0.06</th></th<>	0.06
total 87.35 87.08 87.82 87.52 88.42 87.55 86.38 87.11 87.13 86.87 87.50 87.37 K₂O+Na₂O+CaO 0.00 0.11 0.05 0.09 0.05 0.40 0.18 0.18 0.00 0.12 0.23 0.13 Structural formula b V V V V V V V V V 0.00 0.01	0.00
K₂O+Na₂O+CaO 0.00 0.11 0.05 0.09 0.05 0.40 0.18 0.18 0.00 0.12 0.23 0.13 Structural formula b Si 2.85 2.97 2.81 2.89 2.91 3.10 2.96 2.90 2.96 2.90 2.92 3.04 Ti 0.00 0.01 0.00 0.01	0.01
Structural formula b Si 2.85 2.97 2.81 2.89 2.91 3.10 2.96 2.90 2.96 2.90 2.92 3.04 Ti 0.00 0.01 0.00	86.49
Si 2.85 2.97 2.81 2.89 2.91 3.10 2.96 2.90 2.96 2.90 2.92 3.04 Ti 0.00 0.01 0.00 </th <th>0.07</th>	0.07
Ti 0.00 0.01 0.00 1.04 1.10 1.04 1.10 1.08 0.96 Al ^{IVI} 1.21 1.22 1.23 1.31 1.14 1.28 1.17 1.21 1.18 1.15 1.26 1.25 Cr 0.00 0.00 0.01 0.02 0.05 0.01 0.01 0.00 0.01 1.25 Fe 1.47 1.48 1.57 1.54 1.98 1.94 1.85 2.07 1.63 1.19 1.61 1.48 Mn 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 <	
Al 2.36 2.25 2.42 2.42 2.24 2.18 2.21 2.31 2.23 2.25 2.35 2.22 Aliv 1.15 1.03 1.19 1.11 1.09 0.90 1.04 1.10 1.04 1.10 1.08 0.96 Alv 1.21 1.22 1.23 1.31 1.14 1.28 1.17 1.21 1.18 1.15 1.26 1.25 Cr 0.00 0.00 0.00 0.01 0.02 0.05 0.01 0.01 0.00 0.00 0.02 0.01 Fe 1.47 1.48 1.57 1.54 1.98 1.94 1.85 2.07 1.63 1.19 1.61 1.48 Mn 0.01 0.01 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 0.02 Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.02 0.01 0.00 Ca 0.00 0.01 0.00 0.01 0.01 0.01 0.03 0.01 0.02 0.02 0.00 0.01 0.01 0.01 Na 0.00 0.00 0.00 0.00 0.00 0.00 0.01 0.00 0.00 0.00 0.00 0.00 0.00 Octaedral sum 5.97 5.90 5.98 5.89 5.96 5.79 5.94 5.94 5.93 5.97 5.91 5.85	2.96
Aliv 1.15 1.03 1.19 1.11 1.09 0.90 1.04 1.10 1.04 1.10 1.08 0.96 Alvi 1.21 1.22 1.23 1.31 1.14 1.28 1.17 1.21 1.18 1.15 1.26 1.25 Cr 0.00 0.00 0.00 0.01 0.02 0.05 0.01 0.01 0.00 0.02 0.01 Fe 1.47 1.48 1.57 1.54 1.98 1.94 1.85 2.07 1.63 1.19 1.61 1.48 Mn 0.01 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.02 0.01 0.02 0.01 0.00 0.01 0.02 0.01	0.00
Al ^{VI} 1.21 1.22 1.23 1.31 1.14 1.28 1.17 1.21 1.18 1.15 1.26 1.25 Cr 0.00 0.00 0.00 0.01 0.02 0.05 0.01 0.01 0.00 0.02 0.01 Fe 1.47 1.48 1.57 1.54 1.98 1.94 1.85 2.07 1.63 1.19 1.61 1.48 Mn 0.01 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.02 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 <td>2.19</td>	2.19
Cr 0.00 0.00 0.00 0.01 0.02 0.05 0.01 0.01 0.00 0.00 0.02 0.01 Fe 1.47 1.48 1.57 1.54 1.98 1.94 1.85 2.07 1.63 1.19 1.61 1.48 Mn 0.01 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 <th< th=""><td>1.04</td></th<>	1.04
Fe 1.47 1.48 1.57 1.54 1.98 1.94 1.85 2.07 1.63 1.19 1.61 1.48 Mn 0.01 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.02 0.01 0.00 0.01 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.14
Mn 0.01 0.01 0.01 0.02 0.03 0.03 0.03 0.04 0.02 0.02 0.02 Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.02 0.01 0.00 0.01 0.01 0.00 0.01 0.02 0.01 0.00 0.01 0.01 0.02 0.01 0.00 0.01 0.01 0.02 0.01 0.00 0.01 0.01 0.02 0.01 0.00 0.00 0.01 0.02 0.01 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	0.00
Mg 3.27 3.17 3.14 3.01 2.78 2.43 2.84 2.58 3.07 3.58 2.95 3.07 Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.02 0.01 0.01 0.01 0.01 Na 0.00 0.00 0.00 0.00 0.00 0.01 0.02 0.01 0.00 0.00 0.01 K 0.00 0.00 0.00 0.00 0.00 0.01 0.00 <td>1.20</td>	1.20
Ni 0.00 0.01 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.00 0.01 0.02 0.01 0.01 0.01 0.01 Na 0.00 0.00 0.00 0.00 0.00 0.01 0.02 0.01 0.00 0.00 0.01 K 0.00 0.00 0.00 0.00 0.01 0.01 0.00 0.00 0.00 0.00 0.01 0.02 0.01 0.00 0.00 0.01 K 0.00 0.00 0.00 0.00 0.00 0.01 0.00	0.01
Ca 0.00 0.01 0.00 0.01 0.01 0.03 0.01 0.02 0.00 0.01 0.01 0.01 Na 0.00 0.00 0.00 0.00 0.00 0.01 0.02 0.01 0.00 0.00 0.01 K 0.00 0.00 0.00 0.00 0.00 0.01 0.00 0.00 0.00 0.00 0.00 octaedral sum 5.97 5.90 5.98 5.89 5.96 5.79 5.94 5.93 5.97 5.91 5.85	3.56
Na 0.00 0.00 0.00 0.00 0.00 0.01 0.02 0.01 0.00 0.00 0.01 K 0.00 0.00 0.00 0.00 0.01 0.00 0	0.02
K 0.00 0.00 0.00 0.00 0.00 0.01 0.00 0.	0.01
octaedral sum 5.97 5.90 5.98 5.89 5.96 5.79 5.94 5.93 5.97 5.91 5.85	0.00
	0.00
	5.94
vacancies 0.03 0.10 0.02 0.11 0.04 0.21 0.06 0.06 0.07 0.03 0.09 0.15	0.06
Mg# ° 0.69 0.68 0.67 0.66 0.58 0.56 0.61 0.55 0.65 0.75 0.65 0.67	0.75
R ^{2+d} 4.75 4.66 4.74 4.57 4.79 4.41 4.74 4.70 4.74 4.80 4.59 4.57	4.79
Calculated T (°C)	
Bourdelle et al. (2013) e 422 236 494 245 393 152 284 309 267 415 246 188	300
Tcorrected (>350°C) e 343 367 332 340	

Sample	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c	218c
Clast vs matrix	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast
Analysis #	33 / 1	38 / 1	39 / 1	40 / 1	41 / 1	42 / 1	101/	102 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	103 /	104 /
Chlorite composition a																						
SiO ₂	28.75	28.61	28.38	28.00	28.29	28.44	27.80	28.02	27.66	27.81	28.39	28.56	27.99	28.37	28.54	28.60	28.45	27.75	28.02	28.96	28.54	30.11
TiO ₂	0.14	0.02	0.03	0.67	0.05	0.08	0.04	0.04	0.03	0.00	0.06	0.03	0.07	0.10	0.06	0.14	0.06	0.09	0.17	0.39	0.08	0.02
Al ₂ O ₃	18.97	18.75	19.48	19.58	19.71	19.08	19.25	18.25	19.14	19.10	19.93	18.93	19.01	19.32	19.88	19.38	19.07	19.85	19.33	19.66	19.59	18.50
Cr ₂ O ₃	0.41	0.01	0.00	0.02	0.03	0.03	0.33	0.19	0.29	0.18	0.11	0.13	0.00	0.08	0.00	0.04	0.04	0.17	0.00	0.10	0.09	0.12
FeO	19.01	18.52	17.89	16.44	17.60	19.25	20.10	20.73	19.67	18.92	18.31	19.41	19.11	19.49	19.89	19.64	20.55	18.36	19.79	19.49	20.95	13.56
MnO	0.22	0.26	0.27	0.17	0.26	0.21	0.27	0.31	0.28	0.31	0.28	0.22	0.22	0.30	0.35	0.27	0.18	0.17	0.21	0.26	0.34	0.28
MgO	20.59	20.17	21.41	22.15	21.14	19.97	19.22	19.21	18.95	19.65	19.58	19.23	19.61	18.82	19.43	19.90	18.89	20.27	18.44	19.42	18.63	24.56
NiO	0.17	0.07	0.12	0.06	0.08	0.13	0.09	0.04	0.11	0.13	0.05	0.03	0.09	0.00	0.00	0.16	0.09	0.05	0.05	0.08	0.07	0.14
CaO	0.13	0.05	0.04	0.05	0.06	0.07	0.06	0.05	0.05	0.06	0.09	0.09	0.08	0.06	0.09	0.12	0.10	0.09	0.20	0.25	0.11	0.05
Na₂O	0.02	0.04	0.00	0.01	0.04	0.02	0.09	0.02	0.03	0.04	0.00	0.04	0.00	0.05	0.00	0.00	0.00	0.00	0.05	0.05	0.03	0.03
K₂O	0.01	0.00	0.02	0.01	0.00	0.02	0.00	0.00	0.02	0.01	0.02	0.00	0.02	0.05	0.01	0.02	0.00	0.00	0.00	0.01	0.02	0.00
total	88.41	86.49	87.62	87.16	87.26	87.28	87.24	86.87	86.22	86.20	86.82	86.67	86.22	86.65	88.23	88.26	87.43	86.79	86.26	88.67	88.46	87.36
K₂O+Na₂O+CaO	0.16	0.09	0.05	0.07	0.10	0.10	0.15	0.08	0.10	0.11	0.11	0.13	0.11	0.16	0.10	0.14	0.10	0.09	0.25	0.30	0.17	0.07
Structural formula b																						
Si	2.90	2.94	2.87	2.83	2.87	2.91	2.86	2.91	2.88	2.88	2.90	2.94	2.90	2.92	2.89	2.90	2.92	2.84	2.91	2.91	2.90	2.98
Ti	0.01	0.00	0.00	0.05	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.00	0.01	0.00	0.01	0.01	0.03	0.01	0.00
Al	2.25	2.27	2.32	2.33	2.35	2.30	2.34	2.23	2.35	2.33	2.40	2.30	2.32	2.35	2.37	2.31	2.31	2.40	2.36	2.33	2.35	2.16
Al ^{IV}	1.10	1.06	1.13	1.17	1.13	1.09	1.14	1.09	1.12	1.12	1.10	1.06	1.10	1.08	1.11	1.10	1.08	1.16	1.09	1.09	1.10	1.02
Al ^{vi}	1.15	1.21	1.19	1.16	1.22	1.20	1.20	1.14	1.22	1.21	1.30	1.24	1.22	1.27	1.26	1.21	1.23	1.24	1.27	1.24	1.25	1.14
Cr	0.03	0.00	0.00	0.00	0.00	0.00	0.03	0.02	0.02	0.01	0.01	0.01	0.00	0.01	0.00	0.00	0.00	0.01	0.00	0.01	0.01	0.01
Fe	1.60	1.59	1.51	1.39	1.49	1.64	1.73	1.80	1.71	1.64	1.56	1.67	1.65	1.68	1.68	1.66	1.76	1.57	1.72	1.64	1.78	1.12
Mn	0.02	0.02	0.02	0.01	0.02	0.02	0.02	0.03	0.02	0.03	0.02	0.02	0.02	0.03	0.03	0.02	0.02	0.01	0.02	0.02	0.03	0.02
Mg	3.10	3.09	3.23	3.33	3.19	3.04	2.95	2.97	2.94	3.03	2.98	2.95	3.02	2.89	2.93	3.00	2.89	3.09	2.85	2.91	2.82	3.62
Ni	0.01	0.01	0.01	0.00	0.01	0.01	0.01	0.00	0.01	0.01	0.00	0.00	0.01	0.00	0.00	0.01	0.01	0.00	0.00	0.01	0.01	0.01
Са	0.01	0.01	0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.02	0.03	0.01	0.00
Na	0.00	0.01	0.00	0.00	0.01	0.00	0.02	0.00	0.01	0.01	0.00	0.01	0.00	0.01	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.01
K	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
octaedral sum	5.94	5.93	5.97	5.91	5.95	5.93	5.96	5.97	5.94	5.95	5.89	5.91	5.93	5.89	5.92	5.93	5.92	5.94	5.89	5.87	5.91	5.94
vacancies	0.06	0.07	0.03	0.09	0.05	0.07	0.04	0.03	0.06	0.05	0.11	0.09	0.07	0.11	0.08	0.07	0.08	0.06	0.11	0.13	0.09	0.06
Mg# ^c	0.66	0.66	0.68	0.71	0.68	0.65	0.63	0.62	0.63	0.65	0.66	0.64	0.65	0.63	0.64	0.64	0.62	0.66	0.62	0.64	0.61	0.76
R ^{2+ d}	4.73	4.71	4.77	4.74	4.71	4.72	4.71	4.80	4.68	4.71	4.57	4.64	4.71	4.60	4.65	4.70	4.68	4.68	4.59	4.58	4.64	4.78
Calculated T (°C)																						
Bourdelle et al. (2013) e	322	278	418	373	351	303	371	410	313	334	242	250	305	242	280	299	270	349	248	234	269	285
Tcorrected (>350°C) °			341	323	314		323	338														

Sample	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	155	115	115	115
Clast vs matrix	Clast	Clast	Clast	Clast	Clast	Clast	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	Mat	clast	clast	clast
Analysis #	83 / 1	87 / 1	89 / 1	92 / 1	93 / 1	94 / 1	78 / 1	80 / 1	89 / 1	56 / 1	59 / 1	60 / 1	61/1	73 / 1	74 / 1	78 / 1	79 / 1	25 / 1	28 / 1	33 / 1
Chlorite composition ^a																				
SiO ₂	30.66	28.34	28.78	26.99	26.58	27.52	27.14	27.31	28.80	29.88	27.78	26.30	28.12	26.37	26.72	26.72	26.64	28.14	31.06	30.62
TiO ₂	0.05	0.08	0.06	0.00	0.07	0.02	0.00	0.00	0.10	0.07	0.04	0.03	0.10	0.02	0.00	0.19	0.06	0.04	0.00	0.00
Al ₂ O ₃	19.94	20.83	19.88	19.73	20.45	19.92	20.08	19.97	18.05	17.19	19.13	19.38	19.46	20.03	20.13	20.48	20.67	19.12	16.18	15.94
Cr ₂ O ₃	0.50	0.00	0.00	0.20	0.04	0.06	0.18	0.04	0.02	0.61	0.00	0.00	0.00	0.07	0.02	0.07	0.04	0.05	0.00	0.08
FeO	21.27	17.35	15.81	23.87	26.12	22.39	24.36	22.95	23.72	21.24	25.96	27.42	24.31	28.69	28.29	24.56	26.33	17.71	17.14	16.92
MnO	0.33	0.15	0.12	0.14	0.25	0.21	0.31	0.26	0.20	0.43	0.14	0.20	0.16	0.14	0.21	0.22	0.25	0.23	0.25	0.11
MgO	14.43	20.71	22.77	15.71	15.33	17.69	15.54	16.93	17.11	16.92	14.25	14.09	14.37	12.71	13.27	14.93	13.45	21.41	22.53	22.31
NiO	0.06	0.04	0.03	0.00	0.00	0.12	0.02	0.02	0.00	0.12	0.07	0.00	0.04	0.03	0.05	0.01	0.08	0.09	0.10	0.06
CaO	0.26	0.05	0.08	0.07	0.05	0.03	0.07	0.05	0.11	0.15	0.08	0.04	0.10	0.06	0.08	0.03	0.04	0.12	0.32	0.38
Na₂O	0.05	0.04	0.02	0.06	0.04	0.06	0.09	0.06	0.08	0.12	0.04	0.06	0.09	0.05	0.08	0.04	0.01	0.03	0.06	0.06
K₂O	0.04	0.02	0.00	0.01	0.01	0.00	0.05	0.04	0.02	0.03	0.03	0.04	0.09	0.02	0.01	0.01	0.02	0.03	0.06	0.05
total	87.59	87.62	87.55	86.79	88.95	88.01	87.85	87.62	88.22	86.76	87.52	87.54	86.84	88.19	88.86	87.27	87.59	86.97	87.71	86.54
K ₂ O+Na ₂ O+CaO	0.36	0.12	0.10	0.15	0.10	0.09	0.21	0.14	0.22	0.30	0.14	0.13	0.27	0.13	0.17	0.08	0.07	0.18	0.44	0.49
Structural formula b																				
Si	3.13	2.85	2.87	2.85	2.77	2.84	2.84	2.84	2.98	3.11	2.93	2.81	2.96	2.81	2.82	2.81	2.82	2.87	3.12	3.12
Ti	0.00	0.01	0.00	0.00	0.01	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.01	0.00	0.00	0.02	0.01	0.00	0.00	0.00
Al	2.40	2.47	2.34	2.45	2.51	2.42	2.47	2.45	2.20	2.10	2.38	2.44	2.42	2.51	2.50	2.54	2.58	2.30	1.92	1.91
Al ^{IV}	0.87	1.15	1.13	1.15	1.23	1.16	1.16	1.16	1.02	0.89	1.07	1.19	1.04	1.19	1.18	1.19	1.18	1.13	0.88	0.88
Al ^{vi}	1.53	1.32	1.21	1.30	1.28	1.26	1.31	1.28	1.18	1.21	1.31	1.25	1.38	1.32	1.31	1.35	1.39	1.16	1.04	1.03
Cr	0.04	0.00	0.00	0.02	0.00	0.01	0.01	0.00	0.00	0.05	0.00	0.00	0.00	0.01	0.00	0.01	0.00	0.00	0.00	0.01
Fe	1.81	1.46	1.32	2.11	2.27	1.93	2.13	1.99	2.05	1.85	2.29	2.45	2.14	2.55	2.49	2.16	2.33	1.51	1.44	1.44
Mn	0.03	0.01	0.01	0.01	0.02	0.02	0.03	0.02	0.02	0.04	0.01	0.02	0.01	0.01	0.02	0.02	0.02	0.02	0.02	0.01
Mg	2.19	3.10	3.39	2.47	2.38	2.72	2.42	2.62	2.64	2.62	2.24	2.24	2.26	2.02	2.08	2.34	2.12	3.25	3.37	3.38
Ni	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.01	0.01
Са	0.03	0.01	0.01	0.01	0.01	0.00	0.01	0.01	0.01	0.02	0.01	0.00	0.01	0.01	0.01	0.00	0.00	0.01	0.03	0.04
Na	0.01	0.01	0.00	0.01	0.01	0.01	0.02	0.01	0.02	0.02	0.01	0.01	0.02	0.01	0.02	0.01	0.00	0.01	0.01	0.01
К	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.01
octaedral sum	5.65	5.91	5.95	5.93	5.97	5.95	5.93	5.95	5.92	5.82	5.88	5.98	5.83	5.93	5.94	5.89	5.88	5.98	5.93	5.93
vacancies	0.35	0.09	0.05	0.07	0.03	0.05	0.07	0.05	0.08	0.18	0.12	0.02	0.17	0.07	0.06	0.11	0.12	0.02	0.07	0.07
Mg# c	0.55	0.68	0.72	0.54	0.51	0.58	0.53	0.57	0.56	0.59	0.49	0.48	0.51	0.44	0.46	0.52	0.48	0.68	0.70	0.70
R ^{2+ d}	4.04	4.58	4.72	4.59	4.67	4.68	4.58	4.64	4.71	4.51	4.55	4.71	4.41	4.59	4.60	4.52	4.48	4.79	4.84	4.84
Calculated T (°C)																				
Bourdelle et al. (2013) e	116	307	352	306	509	360	312	335	259	175	223	490	186	321	326	272	251			_
Tcorrected (>350°C) e			315		371	318						365								

Sample	173	173	173	173	173	173	173	173	173	173	173	173	173	173	217	217	217	217	217	217	217	217	217	21
Clast vs matrix	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Clast	Mat	Mat	Mat	Mat	Mat	Clast	Clast	Clast	Clast	Clast	Mat	Mat	Mat	Mat	Ma
Analysis #	42 / 1	74 / 1	97 / 1	98 / 1	99 / 1	106 /	107 /	110/	132 /	79 / 1	84 / 1	102 /	104 /	122 /	93 / 1	94 / 1	101/	102 /	103 /	78 / 1	105 /	107 /	109 /	110
Chlorite composition ^a																								
SiO ₂	32.34	29.47	28.70	28.71	28.64	27.91	28.63	28.67	27.48	27.59	28.87	30.76	31.21	30.67	28.90	27.99	31.48	29.84	28.29	34.92	33.07	28.50	28.11	29.
TiO ₂	0.02	0.05	0.06	0.03	0.06	0.05	0.00	0.09	0.05	0.03	0.03	0.00	0.03	0.17	0.05	0.05	0.06	0.06	0.07	0.00	0.00	0.03	0.04	0.0
Al ₂ O ₃	16.46	17.24	17.57	16.92	18.45	18.34	18.48	17.98	19.10	19.48	18.12	16.51	17.47	19.41	18.04	20.45	17.30	20.94	21.62	13.36	14.48	21.46	21.44	18.
Cr ₂ O ₃	0.09	0.02	0.05	0.03	0.00	0.00	0.00	0.00	0.08	1.38	0.01	0.02	0.01	0.05	0.11	0.09	0.03	0.05	0.00	0.10	0.00	0.04	0.03	0.0
FeO	17.27	17.25	23.81	22.83	22.65	26.14	22.63	18.91	19.66	18.81	20.72	16.35	11.11	11.37	20.26	18.07	11.30	10.08	11.02	15.63	9.26	12.71	12.33	12.
MnO	0.19	0.22	0.26	0.33	0.28	0.24	0.33	0.17	0.14	0.15	0.33	0.07	0.10	0.12	0.31	0.21	0.36	0.48	0.52	0.24	0.21	0.54	0.43	0.3
MgO	20.95	22.00	16.67	17.02	17.61	15.23	17.50	20.53	19.60	19.67	18.04	23.58	26.33	25.69	19.01	20.49	25.96	25.47	24.92	22.35	29.26	24.30	24.08	24.
NiO	0.08	0.21	0.04	0.07	0.09	0.05	0.01	0.07	0.02	0.08	0.12	0.11	0.13	0.14	0.09	0.11	0.11	0.01	0.06	0.28	0.26	0.05	0.07	0.0
CaO	0.05	0.07	0.14	0.14	0.10	0.03	0.08	0.16	0.04	0.04	0.11	0.06	0.05	0.05	0.10	0.08	0.14	0.14	0.12	0.30	0.14	0.17	0.15	0.1
Na₂O	0.03	0.03	0.02	0.04	0.00	0.07	0.04	0.04	0.04	0.00	0.15	0.08	0.05	0.09	0.00	0.00	0.01	0.02	0.02	0.04	0.02	0.03	0.04	0.0
K₂O	0.00	0.03	0.01	0.02	0.02	0.00	0.00	0.03	0.01	0.01	0.02	0.01	0.01	0.05	0.00	0.03	0.00	0.00	0.01	0.06	0.02	0.02	0.00	0.0
total	87.48	86.57	87.33	86.14	87.90	88.06	87.70	86.63	86.23	87.25	86.52	87.54	86.50	87.79	86.85	87.59	86.74	87.08	86.64	87.28	86.72	87.87	86.72	86.
K ₂ O+Na ₂ O+CaO	0.08	0.12	0.17	0.19	0.12	0.10	0.12	0.22	0.10	0.05	0.28	0.15	0.11	0.19	0.10	0.12	0.15	0.16	0.15	0.40	0.17	0.23	0.18	0.1
Structural formula ^b																								
Si	3.24	3.01	3.00	3.04	2.96	2.93	2.96	2.95	2.85	2.83	3.00	3.08	3.07	2.98	2.98	2.83	3.10	2.90	2.79	3.47	3.22	2.80	2.79	2.9
Ti	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.0
Al	1.94	2.07	2.17	2.11	2.24	2.27	2.25	2.18	2.34	2.35	2.22	1.95	2.03	2.22	2.19	2.44	2.01	2.40	2.51	1.57	1.66	2.48	2.51	2.1
Al ^{IV}	0.76	0.99	1.00	0.96	1.04	1.07	1.04	1.05	1.15	1.17	1.00	0.92	0.93	1.02	1.02	1.17	0.90	1.10	1.21	0.53	0.78	1.20	1.21	1.0
Αl ^{vi}	1.18	1.08	1.17	1.14	1.20	1.20	1.21	1.13	1.19	1.18	1.22	1.03	1.10	1.20	1.18	1.27	1.10	1.31	1.30	1.04	0.89	1.28	1.30	1.1
Cr	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.11	0.00	0.00	0.00	0.00	0.01	0.01	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.0
Fe	1.45	1.47	2.08	2.02	1.96	2.30	1.96	1.63	1.71	1.61	1.80	1.37	0.91	0.92	1.75	1.53	0.93	0.82	0.91	1.30	0.75	1.04	1.02	1.0
Mn	0.02	0.02	0.02	0.03	0.02	0.02	0.03	0.01	0.01	0.01	0.03	0.01	0.01	0.01	0.03	0.02	0.03	0.04	0.04	0.02	0.02	0.04	0.04	0.0
Mg	3.13	3.35	2.60	2.68	2.71	2.39	2.70	3.15	3.03	3.00	2.79	3.52	3.86	3.72	2.93	3.09	3.81	3.70	3.66	3.31	4.25	3.56	3.56	3.7
Ni	0.01	0.02	0.00	0.01	0.01	0.00	0.00	0.01	0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.00	0.00	0.02	0.02	0.00	0.01	0.0
Са	0.01	0.01	0.02	0.02	0.01	0.00	0.01	0.02	0.00	0.00	0.01	0.01	0.01	0.00	0.01	0.01	0.01	0.01	0.01	0.03	0.01	0.02	0.02	0.0
Na	0.01	0.01	0.00	0.01	0.00	0.01	0.01	0.01	0.01	0.00	0.03	0.02	0.01	0.02	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.01	0.01	0.0
K	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.0
octaedral sum	5.79	5.95	5.91	5.91	5.91	5.93	5.92	5.95	5.97	5.94	5.90	5.95	5.91	5.90	5.91	5.94	5.89	5.89	5.94	5.75	5.95	5.96	5.95	5.9
vacancies	0.21	0.05	0.09	0.09	0.09	0.07	0.08	0.05	0.03	0.06	0.10	0.05	0.09	0.10	0.09	0.06	0.11	0.11	0.06	0.25	0.05	0.04	0.05	0.0
Mg# ^c	0.68	0.69	0.56	0.57	0.58	0.51	0.58	0.66	0.64	0.65	0.61	0.72	0.81	0.80	0.63	0.67	0.80	0.82	0.80	0.72	0.85	0.77	0.78	0.7
R ^{2+ d}	4.59	4.85	4.71	4.74	4.70	4.71	4.69	4.79	4.76	4.64	4.63	4.90	4.80	4.66	4.71	4.65	4.77	4.56	4.62	4.66	5.04	4.65	4.63	4.8
Calculated T (°C)																								
Bourdelle et al. (2013) e	140	312	243	236	258	284	256	341	475	920	223	295	233	234										
Tcorrected (>350°C) °									361	452														

Table S3. In situ LA-ICP-MS trace element analysis (ppm) of 22 chlorite crystals from silicified samples.

Sample	195	195	195	195	155	155	155	155	155	155	155	218ab	218ab	218ab	218ab	218c	218c	218c	173	173	173
Clast vs matrix	clast	clast	clast	clast	clast	clast	matrix	matrix	clast	clast	matrix	clast	clast	matrix	matrix	clast	clast	matrix	matrix	matrix	matrix
Analysis #	Chl1	Chl2	Chl2-1	Chl3b	Chl1	ChI2	Chl3	Chl3-1	Chl4	Chl5	Chl6	Chl1	ChI2	ChI3	Chl4	Chl1	Chl2	Chl3	Chl1	Chl2	Chl3
Ti (47) ^a	47 b	5289	49	116	188	-	3816	384	144	110	218	248	13	29	28	66	215	bdl	23	25	17
Mn (55)	2837	2720	2383	1985	3340	2319	241	267	2734	2733	525	386	75	93	48	485	1725	45	278	317	281
Cr (53)	41	311	249	108	1022	689	124	92	40	55	231	169	30	69	56	44	746	15	720	1662	1107
Co (59)	34	33	33	40	103	72	bdl	bdl	150	146	36	bdl	bdl	bdl	bdl	30	93	bdl	166	69	84
Ni (60)	183	179	180	202	549	401	28	29	482	604	83	101	25	27	12	110	368	bdl	2639	1330	1689
Cu (63)	19	20	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	38	bdl	280	202	15
Zn (66)	207	172	159	152	225	161	16	14	198	180	40	29	bdl	bdl	bdl	25	108	bdl	1216	865	905
V (51)	172	241	156	134	473	405	45	18	209	164	35	39	bdl	bdl	bdl	66	198	bdl	29	29	26

Analyses carried out at GeoRessources (Nancy, France) with a 193 nm GeoLas Pro ArF Excimer laser (Microlas, Göttingen. Germany) coupled with beam homogenization optics. Analyzed with an Agilent 7500c Quadrupole ICP-MS (Agilent, Santa Clara, USA) equipped with an octopole reaction system with enhanced sensitivity optional lenses (Cs type; Agilent). Internal standard: ²⁸Si, calibrated from the mean value of several chlorite microprobe analyses.

^a Element analyzed (isotope)

^b Concentrations in ppm calibrated against the NIST SRM 610 silica glass reference using values given in Pearce et al. (1997). Absolute concentrations (ppm) calculated from equations in Longerich et al. (1996).

Table S4. Temperature of chlorite formation calculated from composition (R^{2+} and Si) of chlorite in equilibrium with quartz, with the thermometer of Bourdelle et al. (2013) for chlorites from matrix, clasts and both, in silicified samples.

T (°C) in matrix and clasts	195	155	218ab	218c	173
Mean T	250	265	291	294	276
Median T	244	273	303	301	257
Minimum T	156	113	152	234	140
Maximum T	350	492	402	349	452
Standard deviation	49	88	61	36	75
Number of analyses	46	40	36	22	14
T (°C) in matrix	195	155	218ab	218c	173
Mean T	244	275	269		287
Median T	236	272	284		234
Minimum T	156	175	152		223
Maximum T	350	365	340		452
Standard deviation	48	63	64		96
Number of analyses	37	11	9		5
T (°C) in clasts	195	155	218ab	218c	173
Mean T	272	262	298	294	270
Median T	269	273	313	301	258
Minimum T	197	113	194	234	140
Maximum T	332	492	402	349	361
Standard deviation	50	96	59	36	66
Number of analyses	9	29	27	22	9