



HAL
open science

Rare elements enrichment in crustal peraluminous magmas: insights from partial melting experiments

Julie Anne-Sophie Michaud, Michel Pichavant, Arnaud Villaros

► To cite this version:

Julie Anne-Sophie Michaud, Michel Pichavant, Arnaud Villaros. Rare elements enrichment in crustal peraluminous magmas: insights from partial melting experiments. Contributions to Mineralogy and Petrology, 2021, 176 (11), 33 p. 10.1007/s00410-021-01855-9 . insu-03431952

HAL Id: insu-03431952

<https://insu.hal.science/insu-03431952>

Submitted on 17 Nov 2021

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

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Rare elements enrichment in crustal peraluminous magmas: insights from partial melting experiments

Julie Anne-Sophie Michaud¹ · Michel Pichavant¹ · Arnaud Villaros¹

Received: 13 April 2021 / Accepted: 13 October 2021

© The Author(s), under exclusive licence to Springer-Verlag GmbH Germany, part of Springer Nature 2021

Abstract

Experiments were conducted to explore the behavior of Li, Rb, Nb, Sn, Cs, Ta, W during crustal melting and test the anatectic origin of rare metal-bearing peraluminous granites such as rare metal granites (RMGs). The experiments were performed under fluid-absent conditions at 800 and 850 °C, 400 MPa and moderately reducing fO_2 ($\Delta FMQ = -0.5$ to -0.8). Starting materials were cores of several millimetres drilled from two natural rocks, a biotite-rich paragneiss (Pg) and a muscovite-rich orthogneiss (Og) enriched in Li, Be, Sn, Cs, W. Both protoliths produced small melt fractions from 8 to 20% vol. Melt distributions were either homogeneously distributed at grain boundaries in the Pg or preferentially associated with muscovite reaction zones in the Og. In the Pg at 800 °C, melting is mainly fluid present, driven by interstitial water at grain boundaries. At 850 °C, biotite dehydration-melting produces peritectic orthopyroxene, hercynitic spinel, ilmenite and alkali feldspar in addition to melt. In the Og, muscovite dehydration-melting generates melt plus peritectic biotite, hercynitic spinel, ilmenite, Al silicates and alkali feldspar. Experimental glasses are nearly homogeneous, silica rich, peraluminous and leucogranitic and their major element compositions differ only little between the two protoliths. In contrast, the trace element concentrations vary as a consequence of chemical and textural heterogeneities in our starting materials. Compared with source rocks, the Og glasses are enriched in Rb, Nb, Ta, W and depleted in Li, Cs and the Pg are enriched in Li, Rb, Cs, W and depleted in Nb, Ta. Mass-balance calculations indicate that during muscovite dehydration-melting, Li, Cs and Rb partition into the melt; whereas Nb, Ta and W are preferentially incorporated in peritectic phases. Li and Cs also partition toward the melt during biotite dehydration-melting. The partitioning behavior of trace elements during crustal melting is a function of the melting reaction and partition coefficients between melt, residual and peritectic phases. Experimental glasses are similar to peraluminous muscovite granites but fail to reproduce RMG compositions. Alternatives to mica dehydration-melting such as fluid-present and residual source melting emphasize the difficulties with an origin of RMGs by purely anatectic processes. Crystallization differentiation might have to be combined with mica dehydration-melting to explain the distinctive geochemical features of RMGs.

Keywords Melting experiments · Granitic magma · Anatectic origin · Rare element distribution

Introduction

The origin of rare elements concentrations (i.e., Li, Rb, Cs, Nb, Ta, W, Sn) in peraluminous crustal melt remains a controversial issue and there is still a need to investigate

potential enrichment mechanisms. Best examples are highly evolved rare metal granites (RMGs) and their volcanic equivalents (e.g., French Massif Central: Raimbault et al. 1995; Raimbault and Burnol 1998; China: Belkasmi et al. 2000; Zhu et al. 2001; Finland: Haapala and Likkari 2005; Algeria: Kesraoui et al. 2000; Namibia: Schmitt et al. 2000; Nigeria: Kinnaird et al. 1985; Egypt: Renno 1997; Canada: Kontak 1990; Pillet et al. 1992; Russia: Raimbault et al. 1995; Mongolia: Kovalenko et al. 1995). They include peraluminous highly phosphorous RMGs (i.e., PHP, Linnen and Cuney 2005), characterized by high concentrations in fluxing elements (F, B, Li, P) and strong enrichments in rare lithophile elements and metals (Sn, Nb, Ta, Be, Cs, Rb, W,

Communicated by Othmar Müntener.

✉ Julie Anne-Sophie Michaud
julie.michaud@outlook.fr;
j.michaud@mineralogie.uni-hannover.de

¹ Université d'Orléans, CNRS, BRGM, ISTO, UMR 7327, 45071 Orléans, France

U). Studies of volcanic analogs (Kovalenko and Kovalenko 1976; Pichavant et al. 1988a, b; Raimbault and Burnol 1998) have demonstrated the key role of magmatic processes in rare elements concentration. Yet, two models are presently competing on the origin of RMGs. Numerous studies (e.g., Cameron 1949; Jahn and Burnham 1969; Raimbault 1987; Černý 1991, Černý 1992; Raimbault et al. 1995; Breaks et al. 2005; Černý et al. 2005; London 2008; Roda-Robles et al. 2018; Hulsbosch 2019) have proposed that rare element PHP-type magmas result from extreme fractionation of less evolved peraluminous parental magmas (e.g., Shearer et al. 1987; Černý and Ercit 2005; London 2005; Cuney and Barbey 2014; Villaros and Pichavant 2019). However, spatial and chronological relationships between less evolved granites and RMGs are not always consistent with derivation of the latter from the former (e.g., Deveaud et al. 2013; Goodenough et al. 2014; Melleton et al. 2015). This has led to the alternative model of origin of RMGs through partial melting of the crust (i.e., the anatexis model; e.g., Stewart 1978; Christiansen et al. 1986; Manning and Hill 1990; Cuney et al. 1992; Shearer et al. 1992; Nabelek et al. 1992a, b; Falster et al. 1997; Roda et al. 1999; Shaw et al. 2016; Simons et al. 2016; Muller et al. 2017). As an illustration, Nabelek et al. (1992a, b) suggested that the diversity of Black Hill pegmatite geochemistry and mineralogy results from variable melting conditions rather than from different degrees of crystallization differentiation. Although enriched melts such as RMGs represent important rare elements reservoirs, not all rare element deposits are related to such highly evolved igneous bodies. There are numerous examples of a strong spatial relationship between, on the one hand, peraluminous muscovite/two-mica granites and W-deposits and, on the other hand, peraluminous biotite granites and Sn-deposits (e.g., Simons et al. 2016; Harlaux et al. 2018; Hulsbosch 2019; Yuan et al. 2019; Neiva 2002). This suggests that peraluminous crustal melts can have specific rare element signatures and levels of enrichments.

Partial melting is a first-order chemical differentiation process which redistributes elements within the crust. Pressure and temperature conditions, the nature and mineralogy of the source rock and melting mechanisms such as presence or absence of a fluid phase are the main factors controlling crustal anatexis. During the melting reaction, incompatible elements partition into the melt while compatible elements are preferentially trapped in minerals. The solid residuum left upon partial melting (restite, Chappell et al. 1987) comprises different types of phases (Villaros et al. 2018a): (1) refractory phases that are not significantly reacted during the melting process; (2) reactant phases initially present in the source in excess than required by the stoichiometry of the melting reaction and which remain residual; (3) peritectic phases that are produced along with melt if the melting reaction is of peritectic or incongruent type and (4) accessory

phases with trace elements as essential structural constituents (ESC, e.g., zircon for Zr) whose behavior is controlled by their solubility in the melt. Thus, rare elements are redistributed and eventually concentrated in the melt according to the melting behavior of their respective mineral carriers. In most crustal source rocks, micas combine high modal abundances and the ability to concentrate a large number of trace elements (e.g., Bea et al. 1994; Nabelek 1999; Yang and Rivers 2000) although for some elements accessory phases are critical (e.g., Bea and Montero 1999). Therefore, an examination of the melting behavior of micas is necessary to understand how rare elements will distribute during crustal anatexis.

The melting relations of muscovite and biotite have been extensively studied during the 70s and 80s (Brown and Fyfe 1970; Burnham 1979). Both minerals are important sources of water for fluid-absent dehydration-melting which represents an important mechanism for granite magma generation (Thompson 1982; Clemens and Vielzeuf 1987; Le Breton and Thompson 1988; Vielzeuf and Holloway 1988; Vielzeuf and Clemens 1992). However, those studies have focused on phase relations and major element compositions of melts and coexisting minerals (e.g., Le Breton and Thompson 1988; Vielzeuf and Holloway 1988; Montel and Vielzeuf 1994, 1997; Patiño Douce and Harris 1998). Few studies have explored the behavior of trace elements during crustal melting. Rushmer (2001) presented preliminary results on the Sr, Ba and Ce signature of melts formed by dehydration-melting of muscovite. Icenhower and London (1995) studied the partitioning of Li, Ba, Sr, Rb, Cs between muscovite, biotite and peraluminous silicic melt at 200 MPa under H₂O-saturated conditions but high proportions of melt were obtained (i.e., ~ 50 vol% at 700 °C). Recently, the generation of Li-enriched melts was investigated from low degree partial melting experiments of a staurolite-bearing gneiss at 700 MPa, 650–750 °C but no trace element data were reported for the glasses (Konzett et al. 2018). The dependence between the trace element signature of the melt and the stoichiometry of the melting reaction has been theoretically explored for pelitic rocks (Harris and Inger 1992; Harris et al. 1995) and only Ba, Sr and Rb were modeled (Harris and Inger 1992). The behavior of Nb and Ta was tackled by Stepanov and Hermann (2013) who showed that Nb preferentially partitions in the residue than in the melt. Ta-enriched melts can be generated by crustal melting in the presence of residual biotite (Stepanov et al. 2014). For Sn, the variable partitioning behavior between leucosome and restite in migmatites was attributed to the stability of Sn-sequestering minerals during melt generation (Wolf 2018; Wolf et al. 2018).

This paper presents results of partial melting experiments on two natural mica-bearing lithologies as a test of mechanisms of rare element enrichment in granitic melts. We focus

on melting textures and reactions, on the nature of peritectic mineral phases and on experimental melt major and trace element compositions, notably for rare lithophile elements such as Sn, W, Li, Nb, Ta. The data demonstrate production of melts enriched in rare metals under specific conditions. They provide insight on the role of micas in the origin of rare elements magmas and stress the critical importance of the melting reaction and peritectic phases/melt element partitioning.

Experimental approach, strategy and variables

This study is aimed at testing the generation of rare element-enriched granitic melts by low degree partial melting of natural crustal source rocks. Degrees of melting (i) are in the 10–15 vol% range for peraluminous silicic magmas including PHP RMGs (Clemens and Vielzeuf 1987; Le Fort et al. 1987; Pichavant et al. 1988a, b). In contrast, in our early exploratory experiments performed under fluid-present conditions, high melt fractions ($i \geq 0.5$, Villaros et al. 2018b) were generated. Therefore, fluid-absent conditions were subsequently imposed and experimental i were controlled from two parameters, (1) temperature (either 800 or 850 °C) and (2) type of source rock. Pressure was kept constant (400 MPa). Low melt fractions were also targeted because, for incompatible elements (Kd mineral/melt < 1), melt concentrations increase markedly when F drops below 0.1–0.2 (e.g., Hanson 1978), an alternative to extreme fractionation with the crystallization-differentiation model. It was expected that the detection of trace elements would be facilitated in glasses from low F charges given that, in this study, natural undoped starting materials have been used. However, for rare element concentrations to be determined, glass pools must be of sufficient size and this illustrates the necessary trade-off between melt fraction and glass analysis in this study.

Sources of peraluminous magmas include an important fraction of metasediments (Le Fort et al. 1987; Pichavant et al. 1988a, b; Ballouard et al. 2015) but there is also isotopic evidence for a metagneous component (e.g., Turpin et al. 1990). Therefore, one paragneiss (Pg) and one orthogneiss (Og) were selected as representative of the main protoliths involved in peraluminous silicic magma generation. The first advantage is that the behavior of each mica can be followed during partial melting (e.g., Rushmer 2001) since mica modal compositions are contrasted between the two samples (the Pg is biotite rich and contains minor muscovite, whereas the Og is muscovite rich with only minor biotite, see below). The second advantage is that the two protoliths have contrasting whole-rock major and trace element compositions. In particular, the Og is clearly enriched

in lithophile trace elements (Li, Be, Cs, Sn, W) compared to the Pg (Table 1 and see below) and so is representative of a pre-enriched geochemically “fertile” source rock.

Instead of powders, microcores were drilled to preserve original textures, such as presence of metamorphic layering and segregation of micas. Similar microcores have been used previously in several experimental partial melting studies (e.g., Rushmer 2001; Holyoke and Rushmer 2002; Misra et al. 2009; Tumarkina et al. 2011). This approach has the major advantage to closely simulate natural reaction mechanisms and generate textures directly relevant to crustal melting processes. The same approach is followed in this study where we focus on mechanisms of melting reactions and on the associated geochemical implications.

Methods and analytical techniques

Experimental methods

Cores (~7–10 mm length, 2.5 mm diameter; Fig. 1) were drilled in each of the two starting rocks perpendicularly to the foliation and then kept in an oven before use. These were loaded without any additional volatile in Au capsules (15–25 mm length, 2.5 mm diameter slightly enlarged to accommodate the core samples, 0.2 mm wall thickness), subsequently welded shut. The Og and the Pg capsules were reacted together at 400 MPa for a duration of two weeks at 800 and 850 °C. The two experiments were performed in an internally heated pressure vessel operating vertically and pressurized with Ar–H₂ gas mixtures (Scaillet et al. 1995). During the experiments, total pressure was permanently recorded and is known to better than ± 2 MPa. Temperature was continuously measured by three thermocouples with an uncertainty of ± 5 °C. At the end of each experiment, the power to the furnace was switched off and temperature dropped by about 300 °C during the first 5 min. Simultaneously, Ar was pumped inside the vessel to keep pressure approximately constant during quench. Capsules were checked for leaks by weighting and then opened. Experimental charges were cut in their middle parallel to their long axis, embedded in epoxy and then polished.

To generate a moderately reducing environment in the experiments, the proportion of H₂ in the pressurizing gas mixture was precisely adjusted. H₂ was introduced first to a constant P_{H_2} of 0.60–0.61 MPa and then Ar was pumped to a total pressure of ~230–250 MPa before heating up. Experimental f_{H_2} were measured from CoPd solid sensor capsules (Taylor et al. 1992) placed in the vessel together with experimental capsules. After experiments were completed, atomic fractions of Co (X_{Co}) in CoPd alloys were measured (X_{Co} = 0.45–0.46 at 800 °C, 0.39–0.43 at 850 °C) and the sensor f_{O_2} and experimental f_{H_2} (5.1–5.3 MPa at 800 °C,

Table 1 Whole rock compositions of the starting materials

Major elements (wt%)	Pg	Og	Detection limits
SiO ₂	69.29	72.03	0.05
TiO ₂	0.81	0.3	0.02
Al ₂ O ₃	13.72	13.56	0.04
FeO	4.6	1.04	0.015
MnO	0.06	0.02	0.015
MgO	1.76	0.44	0.03
CaO	1.25	1.48	0.03
Na ₂ O	3.4	1.66	0.02
K ₂ O	2.68	5.94	0.03
P ₂ O ₅	0.2	1.25	0.1
F	0.05	0.35	0.002
LOI	0.89	2.05	
Total	99.22	99.89	
A/CNK	1.27	1.14	
A*	57.6	33.1	
B*	117.8	29.0	
Trace elements (ppm)			
Li	48	95	0.5
Be	2.21	6.46	0.05
As	11.98	16.56	0.5
Ba	844.1	745.2	5.5
Bi	0.18	0.2	0.045
Cd	0.35	0.24	0.02
Ce	102.1	66.62	0.03
Co	9.27	1.5	0.08
Cr	109	96	0.5
Cs	6.64	34.5	0.02
Cu	10.73	bdl	2
Dy	5.03	5.74	0.004
Er	2.68	2.95	0.002
Eu	1.44	1.34	0.002
Ga	19.15	22.4	0.02
Gd	5.99	6.07	0.005
Ge	1.97	2.11	0.04
Hf	11.77	5	0.03
Ho	0.95	1.11	0.001
In	0.08	bdl	0.03
La	55.59	37.93	0.02
Lu	0.46	0.39	0.001
Mo	bdl	3.3	0.5
Nb	16.7	11.12	0.015
Nd	43.35	35.52	0.016
Ni	23.42	47.66	2
Pb	20.26	26.87	0.45
Pr	12	9.41	0.004
Rb	110.2	245.7	0.15
Sc	12.04	4.16	0.6
Sb	0.27	2.02	0.06
Sm	7.75	7.66	0.005

Table 1 (continued)

Major elements (wt%)	Pg	Og	Detection limits
Sn	2.81	217.9	0.3
Sr	175.8	98	0.7
Ta	1.78	0.93	0.004
Tb	0.9	0.98	0.001
Th	17.33	14.5	0.015
Tm	0.42	0.43	0.001
U	3.92	2.57	0.01
V	90.88	17.41	0.85
W	2.66	16.96	0.8
Y	27.74	31.44	0.02
Yb	2.87	2.73	0.002
Zn	82.59	25.08	7
Zr	537.1	185.6	1.5
Nb/Ta	9.4	12.0	

bdl below detection limit

$A/CNK = Al_2O_3 / (Na_2O + K_2O + CaO)$ molar

$A^* = 1000 \cdot (Al - (Na + K + 2Ca))$ atomic (Debon and Le Fort 1983)

$B^* = 1000 \cdot (Fetot + Mg + Ti)$ atomic (Debon and Le Fort 1983)

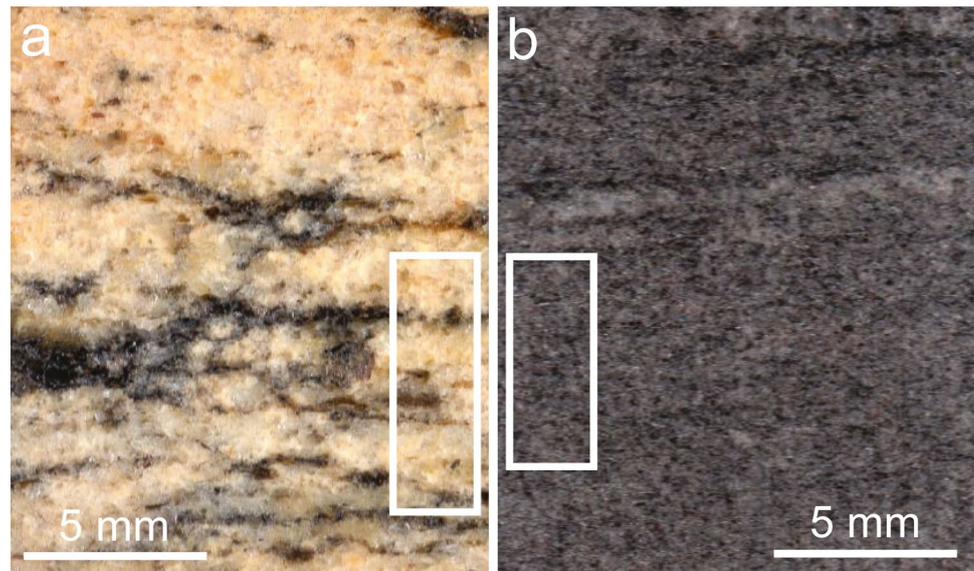
4.7–5.9 MPa at 850 °C) were determined (e.g., Scaillet et al. 1995). The fO_2 within experimental capsules was calculated from the determined fH_2 , the formation constant of water (K_w , Robie et al. 1979) and the fH_2O , the latter obtained assuming concentrations of 4 wt% H₂O in glass using the by-difference method (Villaro et al. 2018b). In this study, fO_2 was similar in the four experimental capsules, yielding deviations from the fayalite–magnetite–quartz oxygen buffer (ΔFMQ) of -0.7 to -0.8 at 800 °C and of -0.5 to -0.7 at 850 °C.

Analytical methods

The two starting rocks were analyzed for major and trace elements at the Centre de Recherches Pétrographiques et Géochimiques (CRPG-SARM, Vandoeuvre les Nancy) by ICP-OES and ICP-MS (see details of the method in Carignan et al. 2001).

Starting and experimentally reacted samples were imaged with a Merlin compact ZEISS scanning electron microscope (SEM, ISTO, Orléans) working under an acceleration voltage of 15 kV and equipped with a Bruker EDS detector. Backscattered electron (BSE) images (panoramas of entire experimental charges and high magnification views of detailed microtextures) and potassium X-ray distribution maps were acquired. The 2D images were processed to calculate experimental melt volumetric proportions. High-resolution BSE panoramas (glass and minerals distinguished from each other by gray level and texture) were superposed

Fig. 1 Macroscopic views of the two starting materials. **a** Og; **b** Pg. Note the differences in deformation intensities, the Pg being more finely banded than the Og, and the heterogeneous mica distribution in the Og. The white rectangles give sizes of experimental microcores



with potassium distribution maps, to image the glassy areas. K_2O is a good tracer for melt pools as concentrations in melts are similar in all experimental charges (i.e., between 5 and 6 wt%), and differ from other minerals such as micas. Melt zones were manually outlined and black and white images produced for the whole sample to count the number of pixels associated to 100% melt areas and the total number of pixels (Fig. A1). Muscovite and biotite reaction zones with melt intimately intermixed with fine-grained peritectic phases (see below) were also delineated and percentages of melt in these zones estimated (Table A1 and Fig. A2). As an example, in the Og sample at 800 °C, two representative types of melt-peritectic phase mixture zones were recognized (i.e., zones a, b; Table A1 and Fig. A2). Both zones were manually outlined using small-scale BSE images. Then, black and white images were created, pixels counted, and melt proportions were calculated from the respective numbers of pixels for melt and peritectic phases, yielding 63 and 25% melt for zones a and b, respectively (Table A1 and Fig. A2). To extend the calculations at the scale of the whole sample, it was assumed that all mica reaction zones are either of type a (63% melt) or b (25% melt). Finally, the degree of melting was calculated as the number of pixels for 100% melt zones plus the number of pixels for melt in all mica reaction zones divided by the number of pixels for the whole sample. Pixel counting was performed using the ImageJ software (e.g., Schneider et al. 2012).

Mineral phases (starting materials and experimental samples) and glasses (experimental samples) were analyzed for major and minor elements using a CAMECA SX Five electron microprobe (ISTO, Orléans) operated under an acceleration voltage and a beam current of 15 kV and 6 nA, respectively. Standards included albite ($SiK\alpha$ and $NaK\alpha$), synthetic $MnTiO_3$ ($MnK\alpha$ and $TiK\alpha$), Al_2O_3 ($AlK\alpha$), Fe_2O_3

($FeK\alpha$), MgO ($MgK\alpha$), andradite ($CaK\alpha$), orthoclase ($KK\alpha$), apatite ($PK\alpha$) and topaz ($FK\alpha$). Mineral phases were analyzed with a focused beam (1–2 μm). For glasses, spot sizes of either 15 or 50 μm were used to minimize the migration of alkali and match the laser spot sizes (see below). Results were checked against analyses of Macusani glasses CH0 and JV2 (Pichavant et al. 1987) used as secondary standards. Glass–peritectic phase mixtures in muscovite reaction zones were also analyzed with the same beam sizes (15 and 50 μm). For glasses, relative analytical errors, determined as standard deviations on multiple analyses on Macusani glass CH0, are: SiO_2 (0.5–1%), Al_2O_3 (1–1.5%), FeO (15–25%), MgO (20–40%), CaO (10–17%), Na_2O (2–3.5%) and K_2O (1–2%). The electron microprobe data were also used in mass-balance calculations. Mineral modes in the two protoliths were calculated by mass-balancing the major element chemistries of the main mineral phases against the bulk (i.e., whole-rock) compositions (see Nabelek 1999). Stoichiometries of melting reactions were determined by mass-balancing the major element compositions of reactants and products within the same experimental charge (see Cesare 2000). Calculations were performed on eight oxides (SiO_2 , TiO_2 , Al_2O_3 , FeO , MgO , CaO , Na_2O , K_2O) simultaneously (i.e., using an input matrix with eight rows) and with a variable number of phases depending on the reaction, following the linear least square procedures detailed in Albarède (1995).

Trace element concentrations were determined by laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) at Orléans (joint IRAMAT-ISTO facility). A RESOLUTION (S-155-E) ArF excimer laser (LA; $\lambda = 193$ nm) coupled to a Thermo Finnigan Element XR (extended range, high-resolution) ICP-MS was used. Ablation was performed at 5 Hz, at an energy of ~ 5 $J.cm^{-2}$. SiO_2 from electron

microprobe analysis was used as an internal standard. Mineral phases from the two starting rocks were analyzed with 50 μm spots. Experimental glasses were analyzed with spots of either 15 or 50 μm size depending on the texture and width of glass zones. Glass plus peritectic phase mixtures from muscovite reaction zones were also analyzed with spots of 15 and 50 μm to constrain compositions of bulk peritectic phase assemblages. NIST610 was used as an external standard (Pearce et al. 1997) and Macusani glasses CHO and JV2 (Pichavant et al. 1987) as control. Reference glasses (NIST610 and Macusani) were analyzed three times at the beginning and at the end of each run of ten analyses. For mineral phases, 39 isotopes (Li^7 , Be^9 , B^{11} , Na^{23} , Al^{27} , Si^{29} , P^{31} , K^{39} , Sc^{45} , Ti^{49} , V^{51} , Zn^{66} , Zn^{68} , Rb^{85} , Sr^{88} , Y^{89} , Zr^{90} , Nb^{93} , Sn^{118} , Cs^{133} , Ba^{137} , La^{139} , Ce^{140} , Pr^{141} , Nd^{144} , Sm^{147} , Eu^{153} , Gd^{157} , Tb^{159} , Dy^{163} , Ho^{165} , Er^{167} , Tm^{169} , Yb^{172} , Lu^{175} , Hf^{178} , Ta^{181} , W^{182} , Pb^{208}) were analyzed and for experimental glasses and glass–peritectic phase mixtures 21 isotopes (Li^7 , Be^9 , B^{11} , Na^{23} , Al^{27} , Si^{29} , P^{31} , K^{39} , Cu^{63} , Zn^{64} , Zn^{66} , Rb^{85} , Sr^{88} , Nb^{93} , Sn^{118} , Cs^{133} , Ba^{137} , Ba^{138} , Ta^{181} , W^{182} , U^{238}). The raw data were processed off-line using the Glitter software where time-resolved signals were examined one by one. Element concentrations were calculated for the initial ~ 30 s of all ablation signals relative to the gas blank (~ 20 s), with a linear interpolation across standard analyses.

Results

Source rocks: mineralogy, composition and trace element distribution

The Og and Pg source rocks are Variscan gneisses belonging to the Lower Gneiss Unit (LGU) and were sampled near Chalus in the Limousin part of northwestern French Massif Central (FMC). The LGU is composed of metagreywackes, metapelites and metarhyolites intruded by Cambrian–early Ordovician granitoids transformed into orthogneisses during the Variscan orogeny (e.g., Faure et al. 2009). The LGU underwent two phases of metamorphism during the Middle Devonian and the Late Devonian–Early Carboniferous and mostly of amphibolite grade (Faure et al. 2009). The two rocks are deformed, the foliation in the Og being less strongly marked than in the finely banded Pg (Fig. 1). Both samples are texturally homogeneous and there is no evidence for migmatization and partial melting. Mass-balance calculations using the whole-rock (Table 1) and the major element compositions of the main mineral phases (Tables 2, A2) show that the Og is remarkably muscovite rich with 35% quartz, 29% alkali feldspar, 18% plagioclase, 3% biotite and 13% muscovite. The rock also contains 2% apatite and a few garnets. Using the same calculations, the Pg hosts 34% quartz, 5% alkali feldspar, 37% plagioclase, 20% biotite, 3% garnet, 1% kyanite and contains minor muscovite. Pressure–temperature estimations with mineral thermobarometers and PerpleX pseudo sections constructed using bulk rock and mineral compositions have yielded conditions of metamorphic equilibration of 600 °C–450 MPa for

Table 2 Representative EMPA compositions of micas in the Pg and Og starting materials

Major elements (wt%)	Pg		Og						
	Bt		Bt		Ms				
SiO_2	35.235	35.729	35.403	33.843	34.536	34.538	48.359	47.877	47.349
TiO_2	3.537	3.992	3.757	2.264	2.473	2.569	0.622	0.511	0.782
Al_2O_3	18.507	18.516	18.922	17.1	16.69	16.875	32.46	32.507	32.477
FeO_{tot}	17.685	17.47	18.032	27.365	26.105	26.285	2.98	3.24	2.968
MnO	0.064	0.128	0.108	0.345	0.303	0.307	0.00	0.106	0.023
MgO	8.027	8.18	7.92	2.913	3.375	3.452	0.725	0.67	0.673
CaO	–	–	–	–	–	0.011	0.025	0.019	–
Na_2O	0.088	0.158	0.187	0.036	0.026	0.08	0.435	0.393	0.362
K_2O	9.55	9.313	9.472	9.135	8.894	9.035	9.594	8.98	9.664
P_2O_5	0.147	–	0.038	0.038	0.008	0.145	–	0.023	0.051
F	0.149	0.162	0.139	0.349	0.4	0.404	0.161	0.267	0.185
Total	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Mg#	0.45	0.45	0.44	0.16	0.19	0.19	0.30	0.27	0.29

Bt biotite, Ms muscovite, FeO_{tot} total Fe as FeO

Mg# = $\text{Mg}/(\text{Mg} + \text{Fe})$ at

the paragneiss and of 650 °C–400 MPa for the orthogneiss (Gaudel 2016).

The major element data (Table 1) show that the Pg is more mafic (FeO = 4.60 wt%, MgO = 1.76 wt%, $B^* = 118$) than the Og (FeO = 1.04 wt%, MgO = 0.44 wt%, $B^* = 29$), whereas the Og is more potassic (5.94 vs. 2.68 wt% K_2O) and remarkably P_2O_5 rich (1.25 wt%). Both rocks are peraluminous, the Pg more than the Og ($A/CNK = 1.27$ and 1.14; $A^* = 58$ and 33, respectively). In terms of trace elements, the Pg is enriched in Ta (1.78 ppm), Nb (16.70 ppm), LREE (Ce = 102.1 ppm, La = 55.59 ppm), Zr (537.1 ppm)

and Th (17.33 ppm) compared to the Og. This latter sample is consistently enriched in rare elements and metals such as Li (95 ppm), Be (6.46 ppm), Cs (34.5 ppm), Sn (217.9 ppm) and W (17.4 ppm) when compared to the Pg.

The trace element data (Table 3) show that, in the Pg, biotites are enriched in Li (~ 163–218 ppm) and contain relatively high amounts of Rb, Nb, Sn, Cs, Ba, Ta and W, although in quite variable concentrations (e.g., ~ 56–126 ppm Nb, ~ 3–12 ppm Ta). The other major phases have lower rare lithophile element and metal concentrations but we note that garnet and quartz contain some Li and plagioclase some Be

Table 3 Representative LA-ICP-MS trace element compositions of micas from starting materials

Trace elements (ppm)	Pg			Og					
	Bt			Bt	Ms				
Li	218.65	188.69	163.22	58.52	178.26	209.07	74.22	87.86	78.73
Be	0.97	0.85	0.92	1.85	2.10	0.95	2.59	2.43	2.47
B	2.99	3.85	5.32	3.84	1.04	1.68	7.40	6.71	7.06
Sc	36.04	36.95	36.34	25.03	19.38	16.90	38.10	42.46	38.54
Ti	21,034.0	21,444.0	21,074.0	5772.7	18,861.4	14,128.0	3815.1	4032.9	4147.4
V	393.6	361.1	349.1	48.72	66.59	69.41	50.77	53.26	57.47
⁶⁶ Zn	346.3	349.9	310.3	28.70	411.5	461.3	32.75	37.39	32.71
⁶⁸ Zn	300.4	317.0	300.2	40.73	206.4	179.1	46.14	50.49	43.52
Rb	488.7	495.9	466.1	558.3	1034.2	1029.0	583.8	682.8	565.4
Sr	1.56	1.49	1.85	2.66	2.81	1.53	5.03	3.74	4.64
Y	0.47	0.23	0.28	0.07	37.64	21.55	0.06	0.11	0.58
Zr	1.69	0.97	1.58	2.68	10.86	2.95	2.70	2.64	3.30
Nb	56.48	126.5	119.0	86.97	227.8	125.9	80.68	81.90	76.64
Sn	11.37	10.68	10.22	48.26	30.53	18.64	66.24	58.74	63.67
Cs	31.14	30.84	29.31	4.54	36.17	36.18	5.45	7.63	5.42
Ba	2133.1	1996.4	1946.7	231.8	162.2	179.4	246.1	284.0	235.9
La	0.14	0.04	0.03	bdl	7.19	3.00	bdl	bdl	0.11
Ce	0.29	0.05	0.10	bdl	6.51	3.11	bdl	bdl	0.11
Pr	0.04	bdl	0.02	bdl	1.98	0.88	bdl	bdl	0.03
Nd	0.14	bdl	bdl	bdl	9.40	4.33	bdl	bdl	0.09
Sm	bdl	bdl	bdl	bdl	2.25	1.13	bdl	bdl	bdl
Eu	0.07	0.05	0.06	0.03	0.38	0.21	0.04	0.04	0.05
Gd	0.13	bdl	0.08	bdl	3.52	1.88	bdl	bdl	bdl
Tb	0.02	bdl	bdl	bdl	0.62	0.35	bdl	bdl	bdl
Dy	0.13	bdl	bdl	bdl	4.28	2.42	bdl	bdl	0.07
Ho	0.03	bdl	bdl	bdl	0.93	0.53	bdl	bdl	bdl
Er	0.10	0.04	0.04	bdl	2.58	1.39	bdl	bdl	0.06
Tm	bdl	bdl	bdl	bdl	0.37	0.17	bdl	bdl	0.01
Yb	0.08	bdl	bdl	0.06	2.38	1.03	0.09	0.11	0.11
Lu	0.02	bdl	bdl	0.02	0.33	0.14	bdl	0.02	0.04
Hf	0.12	0.08	0.07	0.17	0.58	0.21	0.18	0.19	0.29
Ta	3.25	11.73	9.14	2.88	10.33	6.01	2.05	2.54	2.22
W	11.84	11.71	12.23	51.25	4.65	5.34	103.4	68.53	73.84
²⁰⁸ Pb	3.91	3.55	3.58	1.54	3.56	2.08	3.22	2.21	2.73
Nb/Ta	17.37	10.78	13.02	30.22	22.06	20.95	39.33	32.29	34.59

bdl below detection limit, Bt biotite, Ms muscovite

(Table A3). Trace element concentrations in biotite from the paragneiss account for > 90% of the whole-rock Li, Rb, Nb, Cs, Ta, W concentrations and for > 80% of the Sn. In the Og, biotite also contains significant and dispersed concentrations of Li (~ 58–209 ppm), Rb (~ 558–1034 ppm), Nb (~ 87–228 ppm), Sn, Cs Ta (~ 3–10 ppm) and W (Table 3). Muscovite has lower and more homogeneous Li (~ 74–87 ppm), Rb (~ 565–683 ppm), Nb (~ 77–82 ppm), Cs (~ 5–8 ppm), Ta (~ 2–3 ppm) and higher Sn (~ 59–66 ppm) and W (~ 69–103 ppm) than biotite, although, for Li and Rb, concentrations overlap. Concerning the other major phases, K-feldspar contains Rb (423 ppm) and > 1 ppm Li, Nb, Sn, Cs and W, plagioclase Li (17 ppm) and Rb (123 ppm), quartz > 1 ppm Rb and Sn and garnet Li (46 ppm), Rb (94 ppm), Cs (8 ppm) plus > 1 ppm Sn (Table A3).

In the orthogneiss, > 90% of the whole-rock Rb, Nb, W and > 70% of Ta reside in micas. However, the mica Li, Cs and Sn concentrations are globally too low to match the orthogneiss whole-rock analysis. Only 25% of the whole-rock Li and < 10% of the Cs and Sn reside in micas. These deficits cannot be fulfilled by local concentration spikes in muscovite which, despite intensive search, were not found. In the same way, Sn-carriers (e.g., Fe–Ti oxides, titanite) are absent and only a few garnets are present. Thus, the missing trace elements must reside in accessory phases, either inclusions or secondary minerals. It is worth recalling that both the mineralogical and geochemical features of the Og, such as presence of muscovite and peraluminous character, are secondary and derived from metamorphic recrystallization (see Crevola 1987). Garnet commonly appears in FMC orthogneisses developed from early Ordovician granitoids (Lotout et al. 2017). The enriched Og bulk chemistry is interpreted to reflect trace element supply by metamorphic fluids and their trapping in major and accessory phases (e.g., Cuney and Barbey 2014). The accessory phase assemblage probably includes white mica given the unusually high Li, Rb and Cs concentrations analyzed in garnet. Cassiterite microcrystals are also likely to be involved to match the elevated whole-rock Sn content (218 ppm, Table 1).

Partial melting textures and microtextures

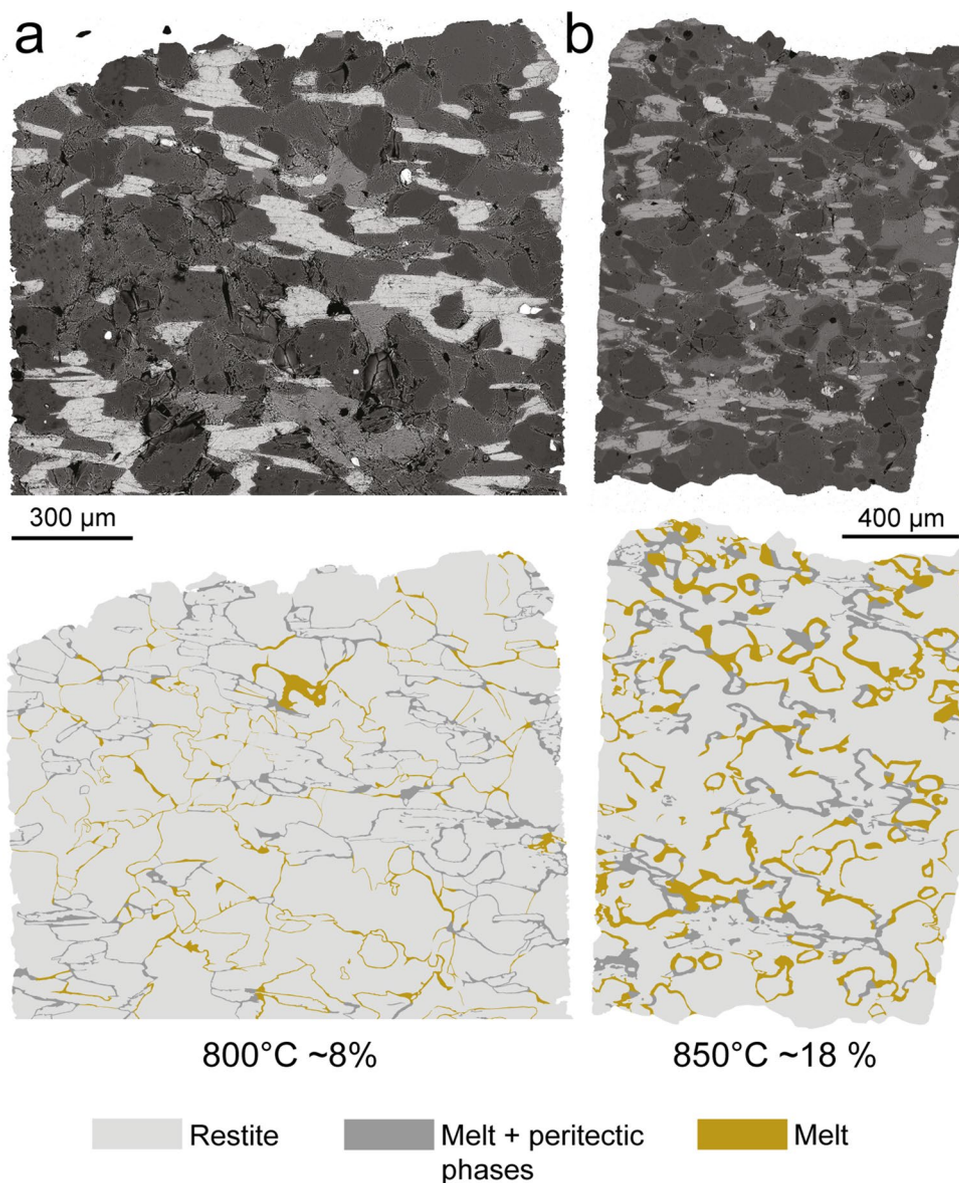
For the two temperatures (800, 850 °C) investigated, both the Og and the Pg underwent partial melting. The two starting rocks yielded quite contrasted melting textures at the millimetre to centimetre scale; whereas, at smaller scales (1–100 µm), microtextures are similar.

Paragneiss. In the Pg changes at 800 and 850 °C, melt forms small lenses and filaments distributed at grain boundaries. Element distribution maps emphasize the homogeneity of melt repartition in the reacted core and the continuity of melt films over distances of several hundreds of µm. The distribution of the melt network is not

influenced by the pre-existing microstructure and foliation. Widths of melt films increase from 800 to 850 °C (Fig. 2) and melt proportions determined from image analysis range from ~ 8 vol% at 800 °C to ~ 18 vol% at 850 °C. At smaller scales, at 800 °C, biotite is largely residual (Bt_{res} in Fig. 3a–c), only showing incipient reaction to glass plus hercynitic spinel (Spl_{per}) and ilmenite (Ilm_{per} , Fig. 3c), whereas the rare muscovites present in the Pg are extensively transformed (Fig. 3a). Muscovite breaks down to melt plus peritectic phases which include biotite (Bt_{per}), Spl_{per} , Ilm_{per} and aluminum silicates, the latter comprising mullite (Mul_{per}) and possibly sillimanite (Sil_{per} , Fig. 3a). Quartz and plagioclase are rounded, generally surrounded by melt and, together with garnet, are interpreted as residual (Qz_{res} , Grt_{res} , Fig. 3b, c). At 850 °C, biotite shows more rounded edges and larger melt pools, distributed along cleavages, than at 800 °C. Biotite show more advanced transformation marked by the appearance of peritectic hercynitic spinel (Spl_{per}), orthopyroxene (Opx_{per}) and ilmenite (Ilm_{per}) distributed around and inside residual biotite (Bt_{res} , Fig. 3d, e). Quartz and plagioclase are residual (Fig. 3d–f) and alkali feldspar occurs both as a residual (Kfs_{res}) and peritectic (Kfs_{per}), forming euhedral overgrowths on residual plagioclase (Fig. 3e). Garnet is more reacted at 850 than at 800 °C and, when close to biotite, can develop overgrowths with inclusions of ilmenite needles suggesting a locally peritectic origin (Grt_{per} , Fig. 3f).

Orthogneiss. At both 800 and 850 °C, melt pools are heterogeneously distributed and preferentially associated with muscovite reaction zones pre-localized in the mica-rich layers (Fig. 4). Melt proportions from image analysis range from ~ 13 vol% at 800 °C to ~ 20 vol% at 850 °C. At smaller scale, at 800 °C, muscovite is entirely consumed while scarce residual biotites are preserved. Muscovite is entirely replaced by glass plus fine-grained peritectic phases (Bt_{per} , Spl_{per} , Ilm_{per} , Mul_{per} , Sil_{per}) and only muscovite “ghosts” can be recognized (Fig. 5a). Peritectic ilmenite is stocky to acicular (< 5 µm) whereas biotite (< 5 µm) and hercynitic spinel (up to 5 µm) are euhedral to subhedral, sillimanite is acicular and, mullite, skeletal (Fig. 5b). Spinel and biotite form chaplets that underline the former muscovite cleavage while sillimanite and mullite are more clustered (Fig. 5d). Euhedral K-feldspar (Kfs_{per}) crystallizes as overgrowths on residual feldspars (Pl_{res} and Kfs_{res}) and residual quartz shows rounded shapes (Qz_{res} , Fig. 5c). At 850 °C, muscovite is totally consumed and biotite undergoes incipient breakdown. Muscovite is replaced by melt plus the same peritectic phases as at 800 °C (Fig. 5e). Residual biotite (Bt_{res}) is associated with peritectic orthopyroxene (Opx_{per}) in addition to hercynitic spinel (Spl_{per}) and ilmenite (Ilm_{per} , Fig. 5f). Orthopyroxenes have sizes from several to tens µm and occur as

Fig. 2 Large-scale textures and melt distributions in Pg charges at **a** 800 °C and **b** 850 °C. The top views are SEM panoramas and the middle views show the corresponding assemblage distributions. Melt proportions (~8% at 800 °C, ~18% at 850 °C) are calculated from the number of pixels for the different assemblages (melt zones: orange, melt + peritectic phases: dark gray, restite: light gray). See text for additional information. Note that both melt and melt + peritectic phases distribute along a continuous network at grain boundaries



elongated stocky crystals growing perpendicular to edges of residual biotite (Fig. 5f).

Experimental phase compositions

Glasses. Representative individual analyses and averages are reported for the four charges (Table 4; Fig. 6). Experimental glasses are relatively homogeneous. However, a minor chemical heterogeneity is apparent, e.g., for Al_2O_3 , the standard deviation of the population of analyses is, in the four glasses, slightly higher than the analytical error (2–2.6% vs. 1–1.5%, see above). This is also true for Na_2O (6.9–16.6% vs. 2–3.5%) and, in some glasses, for SiO_2 , CaO and K_2O (Fig. 6). Migration of alkali especially Na under the electron beam was not corrected for (e.g., Pichavant et al. 1987). Underestimation of Na would artificially increase A/

CNK. However, glasses from the four charges should be impacted uniformly since they differ compositionally very little and, so, the conclusions below should be unaffected. Electron microprobe analyses sum to between 94 and 97 wt% which corresponds to H_2O concentrations of ~4% with the by-difference method (Villaros et al. 2018b).

Glasses from the four charges are all very similar, being SiO_2 rich (74.0–76.3 wt%, Table 4; Fig. 6) and more peraluminous and felsic than the two starting rocks (see Table 1). A/CNK, A^* and B^* extend to higher values in Pg (1.20–1.46, 38–86 and 23–37, respectively) than in Og glasses (1.25–1.36, 46–67 and 18–29, Table 4). The latter are the most K_2O - and Na_2O rich, consistent with the respective compositions of the starting rocks. CaO concentrations are uniform and very low (0.3–0.4 wt%, Table 4). Upon increasing the temperature and the degree of melting, glasses in

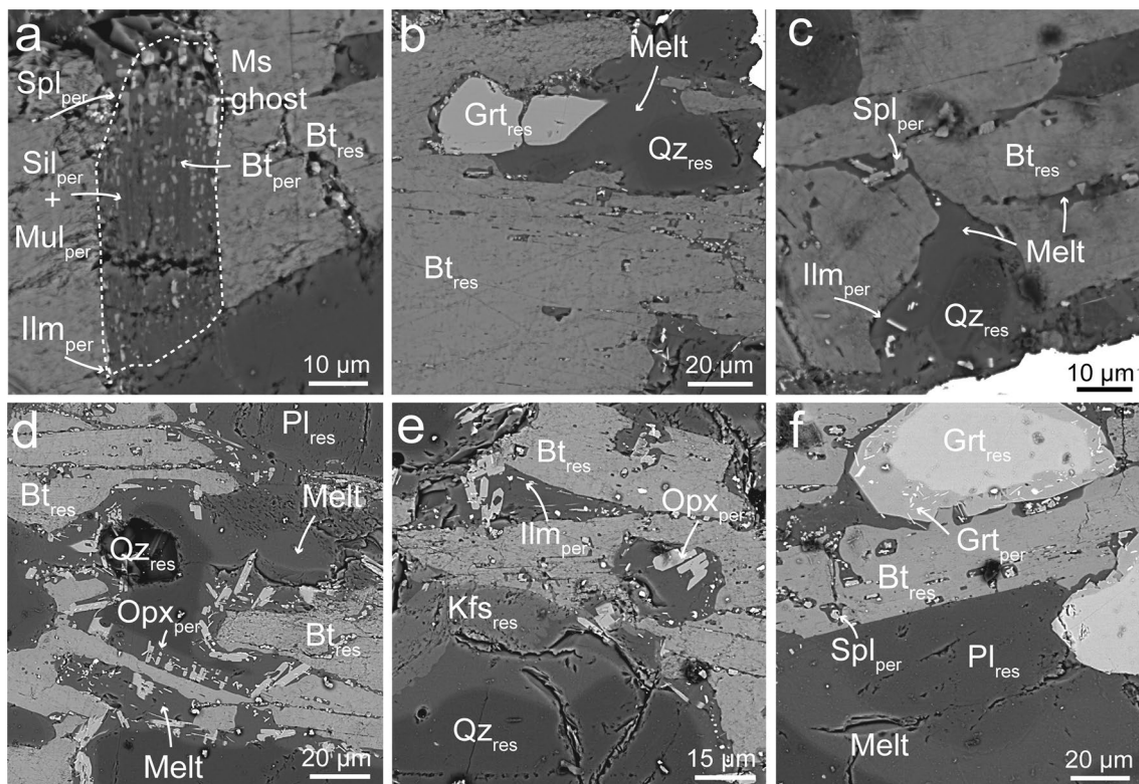


Fig. 3 BSE images showing microtextures in Pg charges at 800 °C (a–c) and 850 °C (d–f). **a** Reacted muscovite “ghost” with melt and peritectic phases including biotite (Bt_{per}), sillimanite and/or mullite (Sil_{per} , Mul_{per}), spinel (Spl_{per}) and ilmenite (Ilm_{per}). **b** Largely unreacted biotite (Bt_{res}) hosting only a few small melt pools with small peritectic phases. Note the rounded edges of residual quartz (Qz_{res})

and the unreacted garnet (Gt_{res}). **(c)** Slightly reacted Bt_{res} and Qz_{res} . Note the presence of Spl_{per} and Ilm_{per} in melt pools. **d, e** Reaction of biotite to form peritectic orthopyroxene (Opx_{per}), Spl_{per} and Ilm_{per} . Feldspars (Kfs_{per} , Pl_{per}) appear mainly residual. **f** Overgrowth of peritectic garnet (Gt_{per}) on residual garnet (Gt_{res}) showing abundant ilmenite inclusions in Gt_{per}

both groups become more K_2O -, CaO - and FeO rich and less peraluminous. Glass FeO/MgO ratios also slightly increase with temperature, consistent with peritectic phases becoming more magnesian at 850 °C (see below). However, these changes are minor, being in the same range or smaller than the standard deviations.

Concentrations of selected trace elements (Li, Rb, Nb, Sn, Cs, Ta, W) in experimental glasses (hereafter designated C_{melt}) are given in Table 5 and the data are illustrated in Fig. 7, where Rb is used as a differentiation index. Most data are for the Og 800 and 850 °C charges and a laser spot size of 15 μm , although, at 850 °C, data with a 50 μm spot were also obtained (Table 5). The raw data were filtered from examination of the 21 measured isotope concentrations (to check glass contamination by mineral phases) and from post-analysis SEM imaging of the charges (to check locations of analytical pits). For the 800 °C Og, only 7 analyses out of 24 spots were retained, and 7 out of 16 for the 850 °C (Table 5). For Pg charges, glass films were generally too thin for LA-ICP-MS analysis and 2 analyses out of 11 spots are available at 850 °C. Both individual and median C_{melt} (instead of mean values

to avoid giving too much weight to outliers) are plotted (Fig. 7). In the 800 °C Og, Li contents range from 62.0 to 103.9 ppm, Rb from 349.2 to 485.0 ppm, Nb from 28.06 to 51.81 ppm, Cs from 5.20 to 19.14 ppm, Ta from 0.43 to 3.89 ppm, W from 23.35 to 52.56 ppm and Nb/Ta from 13.3 to 83.6 (Table 5). At 850 °C, the data are more clustered for some elements (57.5–119.4 ppm Li, 400.1–464.0 ppm Rb, 27.09–44.62 ppm Nb, 10.41–18.56 ppm Cs, 0.55–1.89 ppm Ta, 18.26–37.94 ppm W, Nb/Ta from 21.5 to 45.2, Table 5). Sn is always very low (Og glasses: < 9.33 ppm at 800 °C and < 15.15 ppm at 850 °C; Pg 850 °C glasses: < 2.20 ppm). Glasses from the two protoliths are chemically contrasted as illustrated by the higher Li (167.7–197.6 ppm) in the Pg than in the Og (57.5–119.4 ppm) and the higher Rb (400.1–464.0 ppm) in the Og than in the Pg (250.5–273.1 ppm, data at 850 °C, Table 5; Fig. 7). Ta concentrations are < 2 ppm (except one outlier in the Og at 800 °C) and similar in both glass groups. The Og are globally enriched in Rb, Nb, Ta, W and depleted in Li and Cs compared to the source rock. The Pg are strongly enriched in Li, less so in Rb, Cs and W, whereas both Nb and Ta are depleted. Nb/Ta in the Og (median C_{melt} = 29.0 and 27.4 at

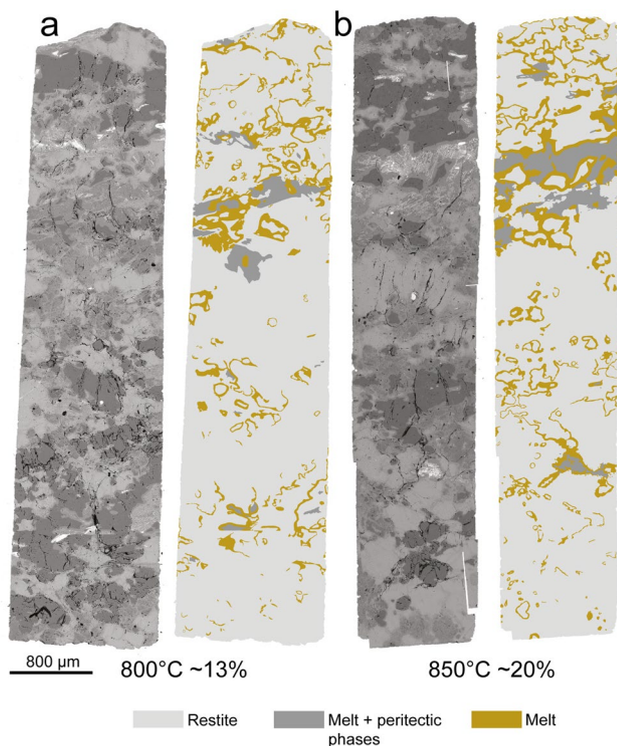


Fig. 4 Large-scale textures and melt distributions in Og charges at **a** 800 °C and **b** 850 °C. For each temperature, SEM panoramas are on the left and assemblage distributions on the right. Calculated melt proportions are ~13% at 800 °C and ~20% at 850 °C. Note that melt (orange zones) and melt + peritectic phases (dark gray zones) are heterogeneously distributed and concentrate in muscovite reaction zones. See text for additional information

800 and 850 °C) and in Pg (13.6, Table 5) glasses are higher than in source rocks (respectively, 12.0 and 9.4, Table 1).

Peritectic phases. Major element compositions of peritectic phases are available in all charges (Tables A4–A6) except the 800 °C Pg because reactions are limited and products, too small for electron microprobe analysis (Fig. 3a, c). Orthopyroxenes (Opx_{per}) at 850 °C are more magnesian ($\text{Mg}\# = \text{at. Mg}/(\text{Mg} + \text{Fe}) = 0.44\text{--}0.53$) in the Pg than in the Og charge ($\text{Mg}\# = 0.19$), consistent with the respective biotite compositions in the protoliths (Table 2), and CaO contents are very low ($< 0.1\%$ Wo). Hercynites (Spl_{per}) also extend to higher $\text{Mg}\#$ in Pg than in Og charges (respectively, 0.34–0.35 vs. 0.22–0.23 at 850 °C) and their $\text{Mg}\#$ decrease with temperature (0.10–0.21 in the Og charge at 800 °C). $\text{Fe}^{3+}/\text{Fe}_{\text{tot}}$ at. proportions in Spl_{per} are mostly between 2 and 9% with no difference between charges. Peritectic biotites (Bt_{per}) in Og charges are more magnesian ($\text{Mg}\# = 0.43\text{--}0.49$) and richer in Al and Ti (Table A5, A6) than in the starting rock whereas residual biotites (Bt_{res}) in the 850 °C Pg charge stay chemically unchanged. Ilmenites (compositions in Og charges only, Ilm_{per} in Pg charges could not be reliably analyzed due to their small size) have $\text{Mg}\# \sim 0.05$ and

a maximum of 3% Fe_2O_3 . Al silicates (Sil_{per} and Mul_{per}) almost exclusively contain Al_2O_3 and SiO_2 but Al/Si at. Ratios are variable from ~ 1 to $> > 1$, i.e., from sillimanite to mullite compositions. Peritectic K-feldspar (Kfs_{per}) is in the range $\text{Or}_{59\text{--}73}\text{Ab}_{26\text{--}37}\text{An}_{2\text{--}4}$, always more Na rich and K poor than in the protoliths (Tables A4–A6). In the 850 °C Pg charge, Kfs_{per} contains ~ 2 wt% BaO (Table A4). Garnet overgrowths in the 850 °C Pg charge (Fig. 3f) are strongly enriched in Mg and depleted in Mn and Ca compared to pre-existing crystals (Tables A2, A4).

The peritectic phases were too small to be analyzed individually for trace elements. Yet, analyses of glass–peritectic phase mixtures from muscovite reaction zones in Og charges constrain the rare element concentrations of peritectic assemblages. A total of 16 analyses are available at 800 °C and 3 at 850 °C (Table 6). Concentrations in mixtures (C_{mix}) are plotted together with C_{melt} for comparison (Fig. 7). Enrichments/depletions in mixtures relative to glass (fractionation factor, $f = C_{\text{mix}}(\text{Table 6})/\text{median } C_{\text{melt}}$ (Table 5)) are quite dispersed because peritectic assemblages in mixtures have variable modal compositions (Fig. 5; Table A7), some being rich in Sil_{per} plus Mul_{per} and other dominated by ferromagnesian phases (Bt_{per} , Spl_{per} , Ilm_{per}) in different proportions. For Rb, f ranges from 0.63 (depleted in mixture) to 1.10 (enriched in mixture), on average 0.90 at 800 °C, slightly higher ($f = 0.99$) and more homogeneous at 850 °C. For Nb and Ta, most f are > 1 (on average ~ 2 at 800 °C, increasing to ~ 3 at 850 °C) indicating enrichments in mixtures. Although scattered (f from 0.99 to 4.74 for Nb and from 0.53 to 5.32 for Ta), the Nb and Ta enrichments are linearly correlated with slopes < 1 (0.3 at 800 and 0.6 at 850 °C) indicating preferential incorporation of Ta over Nb in mixtures. Nb/Ta range widely in mixtures, from 12.2 to 72.0 (800 °C) and 19.9 to 38.3 (850 °C, Table 6), i.e., from lower to higher than glass medians (29.0 and 27.4 respectively, Table 5). Sn concentrations > 20 ppm (Table 6) are found in a few mixtures which contrasts with the low amounts in glasses. W is commonly, although not systematically, enriched in mixtures ($f = 0.71\text{--}5.67$ at 800 °C, 1.18–2.07 at 850 °C). Two W concentrations are very high (above 150 ppm, Table 6) and associated with high Nb and Ta although the Nb–W and Ta–W enrichments are poorly correlated suggesting the presence of a specific W-bearing phase in the peritectic assemblage. Li and Cs have f mostly < 1 and so both are depleted in mixtures compared to glasses. Fractionation factors ranges from 0.11 to 1.43 for Li (on average 0.64 at 800 °C and 0.66 at 850 °C) and Cs from 0.31 to 1.72 (on average 0.75 at 800 °C and 0.79 at 850 °C).

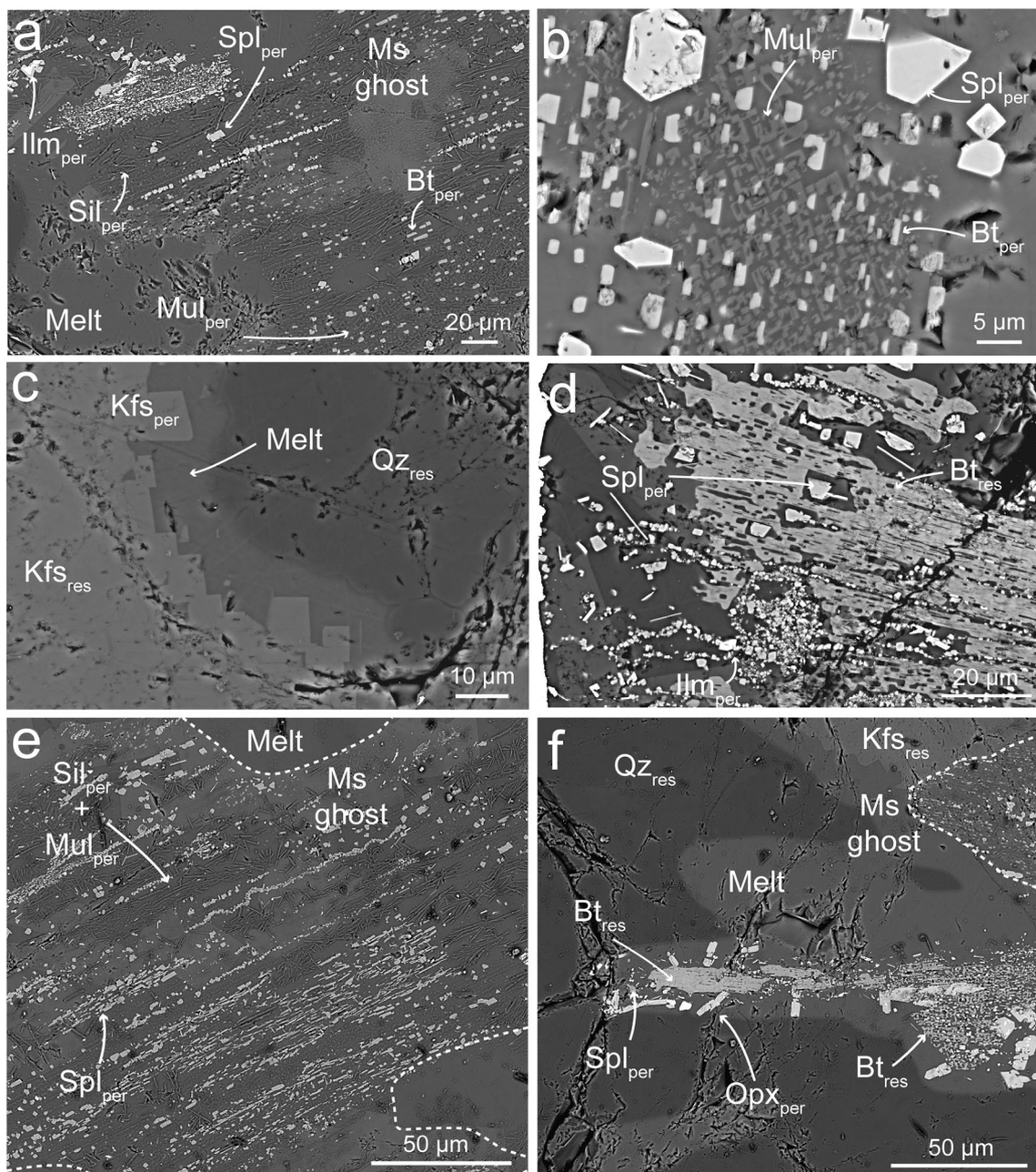


Fig. 5 BSE images showing microtextures in the Og charges at 800 °C (a–d) and 850 °C (e–f). **a** Large-scale view of muscovite reaction zone (“ghost”) showing representative breakdown textures and associated peritectic phases including biotite (Bt_{per}), sillimanite and/or mullite (Sil_{per} Mul_{per}), spinel (Spl_{per}) and ilmenite (Ilm_{per}). **b** Higher magnification view detailing the fine-grained peritectic min-

erals and their textures and the interstitial melt. **c** Peritectic euhedral K-feldspars (Kfs_{per}) overgrowing residual K-feldspars (Kfs_{res}). **d** Biotite showing limited reaction to Spl_{per} and Ilm_{per} . **e** Muscovite reaction zone showing a peritectic assemblage and textures similar to (a). **f** Biotite reacted to peritectic orthopyroxene (Opx_{per}), Spl_{per} and Ilm_{per} . Note the larger melt pools

Discussion

Melting mechanisms and reactions

As emphasized above, melt distribution is contrasted between Bt-rich Pg and Ms-rich Og charges. Differences in

melting behavior between muscovite- and biotite-bearing lithologies have been previously interpreted in terms of contrasted volume changes and dilatational strains associated with the respective mica dehydration-melting reactions (Rushmer 2001; Holyoke and Rushmer 2002). However, textures suggesting melt overpressure (e.g., melt cracks,

Table 4 Representative EMPA compositions of experimental glasses

wt %	P _{Gmelt}						O _{Gmelt}									
	800 °C			850 °C			800 °C			850 °C						
	Median (6)	σ		Median (14)	σ		Median (25)	σ		Median (21)	σ					
SiO ₂	76.27	74.04	75.20	0.92	76.29	74.09	75.43	0.62	76.14	74.75	75.67	0.40	76.27	74.01	75.69	0.60
TiO ₂	0.24	0.25	0.26	0.15	0.26	0.20	0.23	0.05	0.11	0.22	0.15	0.09	0.05	0.24	0.15	0.09
Al ₂ O ₃	13.93	14.74	14.17	0.37	13.58	13.84	13.93	0.31	14.09	14.67	14.06	0.32	13.76	14.65	13.76	0.28
FeO	1.10	1.88	1.49	0.40	1.70	1.90	1.59	0.40	0.99	0.99	1.12	0.30	1.01	1.55	1.27	0.36
MnO	0.03	0.03	0.05	0.02	–	0.08	0.00	0.06	–	0.01	0.01	0.06	–	0.02	0.02	0.05
MgO	0.20	0.26	0.25	0.16	0.22	0.33	0.26	0.06	0.12	0.19	0.12	0.04	0.13	0.17	0.13	0.03
CaO	0.29	0.30	0.29	0.03	0.35	0.43	0.37	0.08	0.31	0.39	0.32	0.08	0.34	0.38	0.34	0.04
Na ₂ O	2.62	2.64	2.63	0.18	2.17	3.22	2.51	0.45	2.15	3.00	2.89	0.26	2.70	3.18	2.70	0.30
K ₂ O	5.21	5.06	5.20	0.10	5.29	5.37	5.31	0.17	6.04	5.75	5.52	0.24	5.75	5.68	5.70	0.19
P ₂ O ₅	0.11	0.79	0.25	0.39	0.14	0.53	0.11	0.16	0.05	0.03	0.03	0.05	–	0.14	0.04	0.09
Total*	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100
A/CNK	1.36	1.46	1.39	0.05	1.41	1.20	1.32	0.10	1.36	1.27	1.27	0.05	1.25	1.25	1.25	0.05
A*	67.23	85.54	72.48	9.01	71.48	37.90	59.18	16.35	67.36	54.46	52.27	10.12	46.36	50.31	47.91	7.91
B*	23.19	35.91	31.94	9.90	32.51	37.27	32.52	6.36	18.10	21.16	21.66	4.04	20.93	28.67	24.11	5.25

*Total normalized to 100%

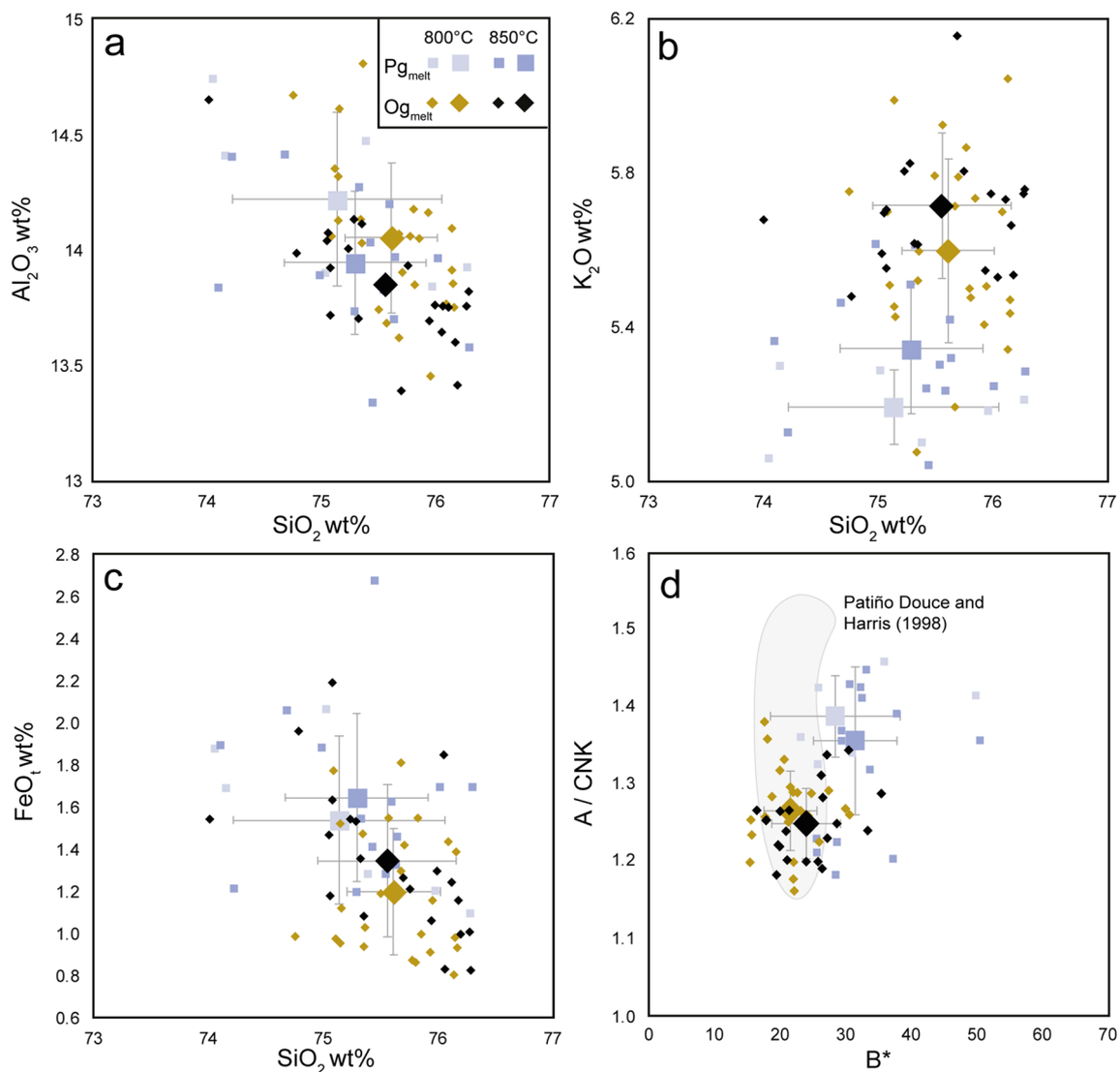


Fig. 6 Major element compositions of experimental melts (paragneiss: Pg_{melt}; orthogneiss: Og_{melt}). Data from Table 4. The small symbols are individual glass analyses (Pg_{melt}: light blue at 800 °C, blue at 850 °C; Og_{melt}: brown at 800 °C, black at 850 °C) and the large symbols represent median values of each of the four populations of analy-

ses. (a)–(c) Hacker diagrams showing Al₂O₃ (a), K₂O (b) and FeO₁ (c) plotted against SiO₂. In (d), data from Patiño Douce and Harris (1998) are shown for comparison. A/CNK is calculated as Al₂O₃/(Na₂O+K₂O+CaO) molar and B* as 1000. (Fe_{tot}+Mg+Ti) atomic as defined in Table 1

Connolly et al. 1997), although locally present, are not widespread in charges from this study and other mechanisms appear to control the melt distribution.

At 400 MPa, 800 °C is well above the muscovite dehydration-melting reaction and in the lower range of the equivalent reaction for biotite (e.g., Spear 1995; Holyoke and Rushmer 2002). 850 °C is in the middle of the range for biotite dehydration-melting (e.g., Spear 1995; Holyoke and Rushmer 2002). Therefore, most muscovite initially present in starting materials would react out at 800 °C; biotite would show only incipient reaction at this temperature and a more advanced transformation at 850 °C. Textural observations confirm this general behavior. At

800 °C, muscovite shows extensive reaction in both Og (Fig. 5a, b, d) and Pg (Fig. 3a) charges. Peritectic biotite is produced, consistent with limited biotite destabilization (Fig. 3c). At 850 °C, clear evidence for advanced biotite breakdown is provided by the appearance of orthopyroxene in both samples (Fig. 3d, e, 5f). However, residual biotite is still present and, so, the “biotite-out” curve is not overstepped (see Holyoke and Rushmer 2002, their Fig. 3).

In the 800 °C Pg charge, the degree of melting is very small (e.g., ~8 vol% from image analysis). Muscovite contributes negligibly to melt formation, being present in minor amount in the starting material. Biotite is abundant, homogeneously distributed, and it remains little reacted (Fig. 3c).

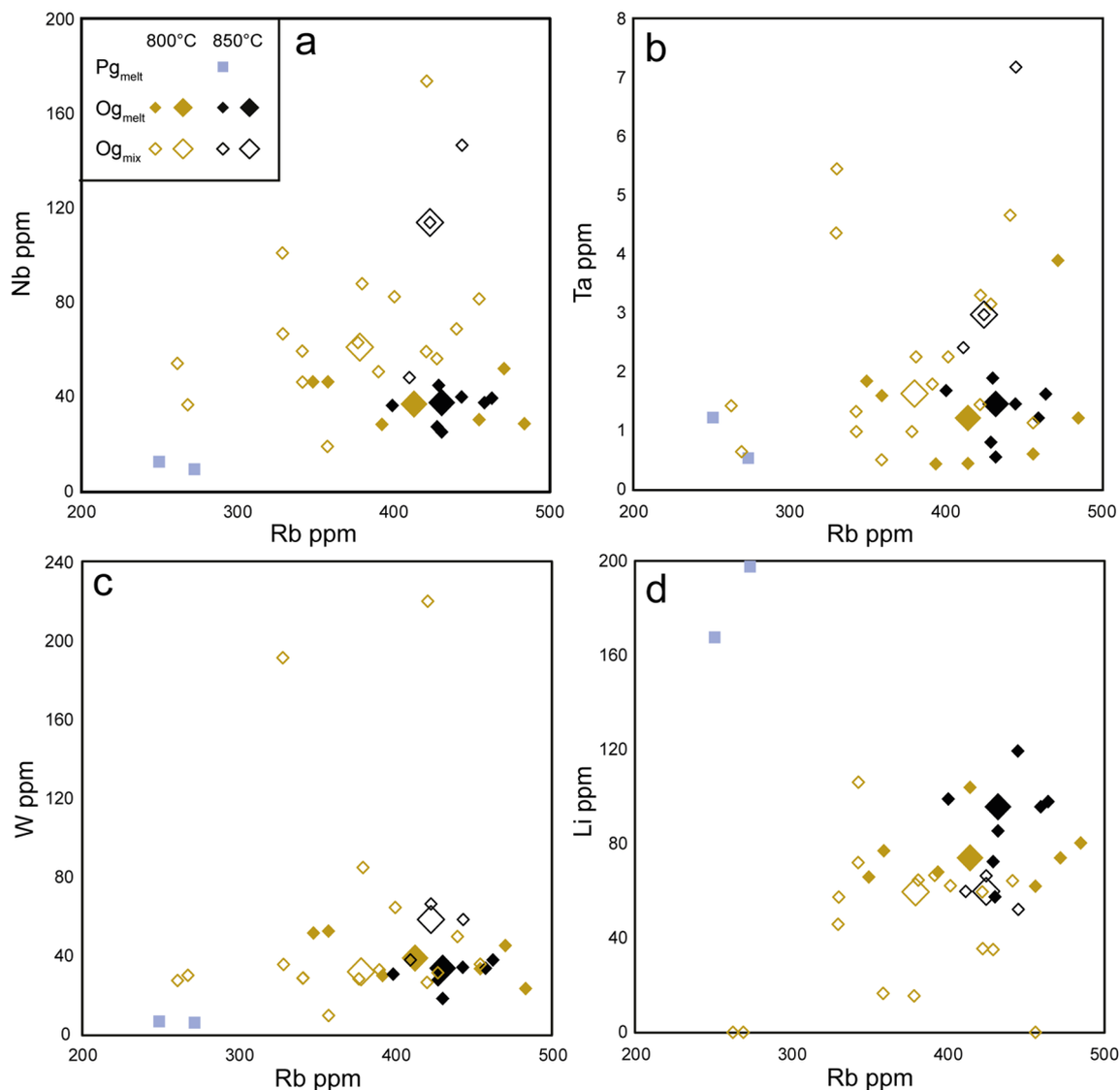
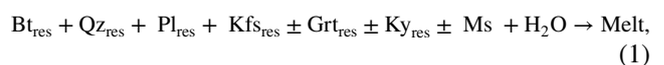


Fig. 7 Trace element compositions of experimental melts (Table 5) and melt-peritectic phases mixtures (Table 6). Only two melt analyses are available for the Pg at 850 °C (small blue symbols, $Pg_{melt}=C_{melt}$). For the Og, the solid symbols are for the melt

($Og_{melt}=C_{melt}$) and the open symbols for the melt-peritectic phases mixtures ($Og_{mix}=C_{mix}$). The small symbols are individual analyses and the large symbols (brown at 800 °C, black at 850 °C) represent median values of each of the four populations of analyses

Biotite edges become rounded, melt pools form along cleavages and peritectic spinel and ilmenite (but not orthopyroxene) appear, the reaction mechanism being similar to the one described by Brearley (1987). Therefore, the contribution of biotite to melt formation is very small. It is recalled that, at 400 MPa, dry melting of a quartz-two feldspar assemblage requires a minimum temperature of ~980 °C (Luth 1969; see also Brearley and Rubie 1990; Johannes and Holtz 1996). Since this temperature is much higher than 800 °C and given that micas are very little reacted, water necessary for melting must be essentially interstitial, present at grain boundaries. Such a water-fluxed mechanism is consistent with the homogeneous melt distribution and the continuous melt

network observed at grain boundaries (Fig. 2). For illustration, assuming 0.1 wt% interstitial water in the Pg (a typical metamorphic pore water concentration, Yardley 2009), ~5.5 vol% melt would be generated at 800 °C, 500 MPa (Clemens and Vielzeuf 1987; Johannes and Holtz 1996), close to estimated (~8 vol% melt). Therefore, the reaction describing melting in the Pg at 800 °C is written as:



where “res” refers to phases which remain residual. Quartz, plagioclase, K-feldspar and biotite (the most abundant phases in the Pg) are the main reactants while garnet, kyanite

Table 6 Concentrations of selected trace elements in glass-peritectic phase mixtures

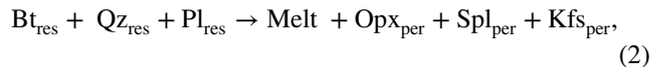
Spot size (μm)	O_{Emix}																					
	850 °C																					
	50	50	50	50	50	15	15	15	15	15	15	15	15	15	15	Median	50	50	50	50	50	Median
Trace elements (ppm)																						
Li	35.11 (1.3)	64.31 (2.3)	64.54 (2.3)	62.19 (2.3)	66.58 (2.5)	16.51 (0.74)	57.37 (10.5)	59.57 (11.5)	72.00 (9.5)	45.82 (7.6)	< 8.26 (4.3)	106.1 (13.7)	15.43 (6.1)	< 14.67 (64.4)	35.53 (7.7)	< 7.74 (4.8)	59.6 (2.3)	66.38 (2.1)	59.75 (2.1)	52.13 (1.8)	59.8 (1.8)	
Rb	428.8 (15.3)	441.3 (16.1)	380.8 (14.1)	401.5 (15.0)	391.3 (15.0)	358.5 (13.5)	330.0 (68.2)	421.9 (31.3)	342.3 (26.2)	329.6 (25.9)	262.2 (21.1)	342.5 (28.4)	378.2 (32.2)	456.0 (40.0)	422.2 (38.2)	268.8 (25.2)	379.5 (14.6)	424.4 (1.8)	411.2 (14.3)	445.0 (15.7)	424.4 (15.7)	
Nb	56.04 (2.2)	68.62 (2.7)	87.78 (3.6)	82.3 (3.4)	50.52 (2.2)	19.17 (0.82)	66.49 (14.8)	59.00 (4.6)	59.26 (5.5)	100.8 (9.5)	53.98 (5.3)	46.02 (4.6)	62.83 (6.5)	81.35 (8.6)	173.9 (18.8)	36.45 (4.2)	61 (4.2)	113.8 (4.2)	47.92 (1.8)	146.6 (5.6)	113.9 (5.6)	
Sn	2.03 (0.2)	2.33 (0.2)	12.36 (0.7)	11.33 (0.6)	9.77 (0.6)	4.96 (0.31)	2.48 (1.0)	7.16 (1.7)	7.98 (1.5)	6.18 (1.5)	7.1 (1.2)	20.87 (2.5)	26.84 (3.1)	23.85 (3.0)	29.40 (3.5)	12.29 (1.8)	8.9 (0.9)	18.75 (0.9)	9.28 (0.5)	6.35 (0.4)	9.3 (0.4)	
Cs	5.33 (0.3)	13.24 (0.6)	6.25 (0.4)	5.96 (0.3)	6.14 (0.4)	1.40 (0.18)	7.14 (1.9)	< 4.87 (2.2)	8.11 (1.8)	4.13 (1.8)	< 2.56 (1.3)	< 4.37 (1.7)	< 4.27 (1.8)	< 4.54 (1.9)	7.19 (2.0)	< 2.38 (1.3)	6.2 (0.5)	10.98 (0.5)	10.91 (0.5)	9.86 (0.5)	10.9 (0.5)	
Ta	3.15 (0.2)	4.66 (0.2)	2.25 (0.1)	2.25 (0.1)	1.79 (0.1)	0.49 (0.04)	5.45 (1.3)	1.44 (0.2)	1.32 (0.2)	4.36 (0.5)	1.42 (0.2)	0.98 (0.2)	0.98 (0.2)	1.13 (0.2)	3.30 (0.4)	0.64 (0.1)	1.6 (0.2)	2.97 (0.2)	2.41 (0.1)	7.18 (0.4)	3.00 (0.4)	
W	31.42 (1.4)	49.84 (2.2)	84.88 (3.7)	64.55 (2.9)	32.89 (1.6)	10.02 (0.52)	35.63 (7.3)	26.44 (2.1)	28.59 (2.8)	191.7 (17.1)	27.43 (2.9)	28.74 (3.0)	28.29 (3.1)	35.73 (4.0)	220.4 (22.5)	30.03 (3.6)	32.2 (2.7)	66.41 (2.7)	37.87 (1.6)	58.47 (2.4)	58.5 (2.4)	
Nb/ Ta	17.8	14.7	39	36.6	28.2	38.9	12.2	41	44.9	23.1	38	47	64.1	72	52.7	57	38.9	38.3	19.9	20.4	20.4	

Data are individual analyses with standard deviations (1 σ) calculated with the Glitter software

and muscovite are accessories. The melt product (composition in Table 4) was mass-balanced ($\Sigma r^2 = 0.08$, Table 7) by a linear combination of the four main reactants (compositions in Table A2), thus validating the stoichiometry of reaction (1). Quartz, plagioclase and K-feldspar are all involved as reactants in approximately equal amounts (~0.3 g per 1 g of formed melt) whereas biotite participates in a smaller proportion (0.13 g per 1 g of formed melt, Table 7), both features being characteristic of water-fluxed melting reactions (e.g. Harris and Inger 1992; Johannes and Holtz 1996). Calculations with garnet and kyanite included as reactants did not change the results significantly and confirm their accessory role in (1). It should be noted that reaction (1) is congruent and must be accompanied by subordinate incongruent reactions (muscovite and biotite dehydration-melting, see below) to account for minor amounts of peritectic phases (Spl_{per} and Ilm_{per}).

At 850 °C in the Pg, the melt distribution is similar to observed at 800 °C. However, the degree of melting is more than twice higher (~18 vol%) than at 800 °C (~8%). The

scarce muscovites initially present are entirely consumed as at 800 °C. The peritectic assemblage changes from 800 (Brearley 1987) to 850 °C and, overall, the biotite dehydration-melting reaction is more advanced. Yet, residual biotite persists in significant amount indicating that the reaction is not overstepped. It can be written as:



where “res” is as above and “per” refers to a peritectic phase. Reaction (2) is closely analogous to the fluid-absent biotite dehydration-melting reaction previously studied in meta-greywackes (Montel and Vielzeuf 1994, 1997) as well as in biotite gneiss and two-micas pelite (Rushmer 2001 and additional references therein). In (2), accessory reactants such as garnet, kyanite and muscovite have been omitted. Yet, garnet is locally involved as illustrated by the development of a Grt_{per} rim on a Grt_{res} near reacting biotite (Fig. 3f). The stoichiometry of reaction (2) was tested by mass-balancing the composition of reactants (Table A2) against the composition

Table 7 Stoichiometries of melting reactions determined by mass-balance

Reaction (1): fluid-present melting, Pg 800 °C									
Reactants				Products					Σr^2
Bt_{res} (g)	Qtz_{res} (g)	Pl_{res} (g)	Kfs_{res} (g)	Melt (g)					
0.13	0.29	0.31	0.31	1					0.08
Reaction (2): biotite dehydration-melting, Pg 850 °C									
Reactants				Products					Σr^2
Bt_{res} (g)	Qtz_{res} (g)	Pl_{res} (g)	Kfs_{res} (g)	Melt (g)	Opx_{per} (g)	Spl_{per} (g)	Ilm_{per} (g)	Kfs_{per} (g)	
1	0.20	0.71	nd	1.15	0.22	0.21	nd	0.07	0.11
Reaction (3): muscovite dehydration-melting, Og 800 °C									
Reactants			Products						Σr^2
Ms (g)	Qtz_{res} (g)	Pl_{res} (g)	Melt (g)	Bt_{per} (g)	Spl_{per} (g)	Ilm_{per} (g)	Als_{per} (g)	Kfs_{per} (g)	
1	0.30	0.43	1.12	0.01	0.03	0.00	0.24	0.28	0.15
Reaction (3): muscovite dehydration-melting, Og 850 °C									
Reactants			Products						Σr^2
Ms (g)	Qtz_{res} (g)	Pl_{res} (g)	Melt (g)	Bt_{per} (g)	Spl_{per} (g)	Ilm_{per} (g)	Als_{per} (g)	Kfs_{per} (g)	
1	0.36	0.34	1.14	0.06	0.01	0.00	0.23	0.26	0.66

Reaction (1) normalized to 1 g of melt; reactions (2) and (3) normalized to 1 g of biotite and muscovite, respectively

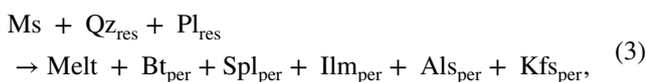
Bt biotite, *Qtz* quartz, *Pl* plagioclase, *Kfs* alkali feldspar, *Opx* orthopyroxene, *Spl* hercynitic spinel, *Ilm*: ilmenite, *Ms* muscovite, *Als* sillimanite-mullite mixture (estimated composition 20 wt% SiO₂, 80 wt% Al₂O₃)

res refers to a residual and *per* to a peritectic phase (see text)

nd phase not considered in the calculations

of products (melt, Table 4; peritectic phases, Table A4) in the Pg 850 °C charge. The low Σr^2 (0.11, Table 7) indicates mass conservation between the left- and right-hand sides of (2). A mass of 1.15 g of melt is produced for 1 g of reacted biotite (Table 7) or $\sim 1.5 \text{ cm}^3$ of melt for 1 cm^3 of biotite. This production rate is broadly consistent with the calculated degree of melting ($\sim 18 \text{ vol}\%$), of which $\sim 8 \text{ vol}\%$ should be accounted for by interstitial water since (2) occurs together with (1) and, so, the degree of melting is controlled by the sum of interstitial water and water released from biotite breakdown. For ~ 10 (i.e., 18–8) vol% melt to be generated with reaction (2), $\sim 7 \text{ vol}\%$ biotite (35% of the initial fraction) must be consumed, consistent with the persistence of a large amount of residual biotite in the charge (Fig. 3d–f).

In the Og 800 °C charge, muscovite has totally reacted out and is replaced by the in situ crystallization of peritectic phases outlining contours of the pre-existing mica (Fig. 5a) and which include Bt_{per} (more Ti- and Mg rich than the starting composition), Spl_{per} , Ilm_{per} , Sill_{per} , Mul_{per} and Kfs_{per} (more Na- and less K rich than the starting composition). This indicates that, as observed in the equivalent Pg charge, the muscovite dehydration-melting reaction (e.g., Spear 1995) has been overstepped. Given the abundance of muscovite in the Og (13 wt%), muscovite dehydration-melting is a major melting mechanism at 800 °C. Melt pools and melt plus peritectic phase mixtures follow the heterogeneous initial distribution of muscovite in the starting microcore (Figs. 1, 4). Biotite is present both as a residual (incipient biotite breakdown results in the formation of small melt pools plus Spl_{per} and Ilm_{per} , e.g., Brearley 1987) and peritectic (Fig. 5b) and, so, the biotite dehydration-melting reaction has not been overstepped, as observed in the corresponding Pg charge. In mica-free zones, quartz and feldspars have melt present at grain boundaries. This could result from muscovite reaction zones being close to but out of the section plane or, alternatively, from interstitial water promoting melting in mica-free zones. However, large amount of interstitial water would induce global melting of the microcore, which is not observed. The muscovite dehydration-melting reaction in the Og at 800 °C can be written as:



where the suffixes “res” and “per” have the same significances as in (1) and (2) and Als_{per} refers to a mixture of sillimanite and mullite. Accessory reactants such as Bt_{res} have been omitted in (3). Reaction (3) is analogous to the muscovite dehydration-melting reactions previously identified in quartzites and pelites (Brearley and Rubie 1990; Connolly et al. 1997; Patiño Douce and Harris 1998; Rushmer 2001). The stoichiometry of reaction (3) has been determined by mass-balancing the compositions of reactants (Table A2)

and melt compositions (Table 4) plus peritectic phases (Table A5). The low Σr^2 (0.15, Table 7) indicates satisfactory mass-balance between the left- and right-hand sides of (3). Kfs_{per} and Als_{per} are the most abundant product phases. Bt_{per} , Spl_{per} and Ilm_{per} are all present in very small amounts ($< 0.05 \text{ g per 1 g}$ of reacted muscovite) consistent with the small concentrations of Fe, Mg and Ti in muscovite. A mass of 1.12 g melt is produced for 1 g of reacted muscovite ($\sim 1.3 \text{ cm}^3$ melt for 1 cm^3 muscovite, Table 7). Given that the Og contains 13 wt% (corresponding to $\sim 13 \text{ vol}\%$) muscovite, total consumption of muscovite following reaction (3) would produce $\sim 16 \text{ vol}\%$ melt, close to but higher than the 13 vol% melt determined from image analysis especially if additional sources of water (mainly interstitial water) are considered. The higher melt fraction in the Og than in the Pg charge at 800 °C (e.g., ~ 13 vs. $\sim 8 \text{ vol}\%$ from image analysis) reflects the higher proportion of muscovite having reacted out in the former than in the latter protolith.

At 850 °C, muscovite dehydration-melting is the essential melting mechanism in the Og, as at 800 °C. Muscovite is totally consumed at the two temperatures and biotite is significantly more reacted at 850 °C than at 800 °C. The formation of peritectic orthopyroxene in addition to ilmenite and spinel (Fig. 5f) marks the same change in peritectic assemblage as already noted from 800 to 850 °C in Pg charges and suggests an increased role of biotite breakdown in melt production, despite a low modal abundance in the Og. However, compositions of melt (Table 4) and peritectic phases (Table 6) differ only marginally between the two Og charges and, as a consequence, the calculated stoichiometry of reaction (3) is little modified from 800 to 850 °C (Table 7). Yet, the quality of the mass-balance calculations is less good at 850 °C than at 800 °C ($\Sigma r^2 = 0.66$ and 0.15 respectively, Table 7). The melt production calculated for 1 g of reacted muscovite is only slightly higher at 850 °C (1.14 g) than at 800 °C (1.12 g, Table 7), which corresponds to a degree of melting of 17 vol% vs. $\sim 20 \text{ vol}\%$ calculated from image analysis in the 850 °C Og charge. In comparison, an increase of 7 vol% in melt fraction (precisely what is observed from image analysis from 13 vol% at 800 °C to 20 vol% at 850 °C) is expected from melt productivity models (Clemens and Vielzeuf 1987; Johannes and Holtz 1996) when temperature is increased from 800 to 850 °C at constant pressure and amount of water. Such an increase in the degree of melting is not captured by our mass-balance calculations. We conclude that, in the Og at 850 °C, the melting reaction is mainly controlled by muscovite dehydration-melting with only minor contribution of biotite and possibly interstitial water to the water budget.

In summary, the two source rocks experienced closely similar melting reactions. Such similarities in melting relations were already stated for hydrous systems by Vielzeuf and Schmidt (2001). At 800 °C, the same muscovite

dehydration-melting reaction (3) takes place in both protoliths and biotite follows the same type of incipient breakdown. Muscovite dehydration-melting is the dominant melt-producing reaction in the Og but has a negligible importance in the Pg which is nearly muscovite free. At 850 °C, biotite breakdown follows reaction (2) both in the Pg and the Og. Biotite dehydration-melting dominates the melting process in the Pg, and, in the Og, minor additional water is supplied. Last, interstitial water contributes to the global melting process following reaction (1), under conditions inappropriate for mica dehydration-melting (Pg at 800 °C) and in combination with the dehydration-melting reactions at 800 and 850 °C. We conclude that both (i) the nature (and the composition) of micas and (ii) the temperature govern melt generation by dehydration-melting. Additional factors such as the presence of interstitial water, the modal abundance of micas and their repartitions in source rocks determine the total melt production and its distribution.

Major element compositions of experimental melts

Major element compositions of melts in this study are tightly grouped (Fig. 6; Table 4) being silica rich, peraluminous and leucocratic, resembling melt compositions from other melting experiments of natural two-mica protoliths generated over a wide range of experimental pressures, temperatures and melt fractions (Patiño Douce and Harris 1998; Rushmer, 2001). Melt compositions were found not to largely vary with the source rock (Patiño Douce and Harris 1998) which is consistent with results from this study as melts from Pg and Og charges differ only marginally (Fig. 6; Table 4). In addition, the Pg and the Og are significantly different (see for example bulk compositions and mica modal abundances), yet Og- and Pg-derived melts have the same major element characteristics than melts generated in other studies (Patiño Douce and Harris 1998; Rushmer 2001).

Melt production in this and previous experimental melting studies is governed by several reactions such as (1)–(3) detailed above (see also Brearley and Rubie 1990; Connolly et al. 1997; Patiño Douce and Harris 1998; Rushmer 2001). Though, the major element compositions of the formed melts are not strongly dependent on the dominant melting reaction. For example, the congruent water-fluxed melting reaction (1) produces a SiO₂-rich, strongly peraluminous and felsic melt (800 °C Pg charge), not significantly different (only slightly more Al₂O₃ rich and FeO_t poor) than the one generated by the biotite dehydration-melting reaction (2) at 850 °C (Table 4; Fig. 6). Melts derived from the muscovite and biotite dehydration-melting reactions (respectively (3) and (2)) differ only marginally. The Og melts are slightly less mafic (i.e., lower B*) and less strongly peraluminous than Pg melts although both are leucocratic (Table 4; Fig. 6). Last, there is no clear indication from the 800 and 850 °C Og

experiments for a significant influence of the melt fraction (from 13 to 20 vol% range according to image analysis) on melt composition. Therefore, major element melt compositions in our experiments are buffered.

One possible explanation is that kinetic factors are important enough to mask the role of melting reactions. Indeed, attainment of equilibrium is an issue in melting experiments of natural polymineralic assemblages at relatively low temperatures, whether single crystals, cores or powders are used (Brearley and Rubie 1990; Patiño Douce and Harris 1998; Hammouda and Pichavant 1999; Rushmer 2001). However, melts analyzed in different parts of the same charge (for example in muscovite reaction zones and in mica-free areas of Og charges) do not show systematic differences. Major element heterogeneities in melts are minor (see above) but their persistence is consistent with melting being kinetically controlled by dissolution of reactant phases and diffusive mass transport within the melt (Hammouda and Pichavant 1999, 2000; Devineau et al. 2005). However, their small magnitude indicates that close-to-equilibrium major element compositions have been reached. This is consistent with the observation that micas at 800 and 850 °C react as expected from equilibrium model reactions (Spear 1995; Holyoke and Rushmer 2002). The reactivity of reactant phases is demonstrated by the compositions of peritectic biotite and K-feldspar which markedly differ from their respective starting compositions and evolve consistently with protolith chemistries (same for peritectic orthopyroxene). Last, the convergence of melt compositions across studies and protoliths (see above) and the uniformity of melt major element chemistries over a large range of pressures, temperatures and melt fractions (Patiño Douce and Harris 1998) are unlikely to be controlled by kinetic factors only because differences in experimental methods and run durations should show up. Therefore, we conclude that near equilibrium major element distributions were attained in our experiments.

As previously emphasized, the same melting reactions take place in both protoliths; water-fluxed and mica dehydration-melting reactions occur simultaneously in the same charge. We note that the mica dehydration-melting reactions (2) and (3) both have spinel, ilmenite and alkali feldspar as products implying that the melts formed are saturated with identical peritectic assemblages, Al silicates and orthopyroxene being excepted. This explains the convergence of melt compositions produced with the two reactions although, in detail, major element compositions of peritectic phases slightly differ between Og and Pg and so melt compositions are not identical. The lack of peritectic Al silicates in reaction (2) is made up for by the presence of accessory kyanite in Pg which imparts a strong peraluminous character to the produced melt, a mechanism which also applies to reaction (1). The appearance of orthopyroxene at 850 °C is not associated with a systematic change in melt chemistry. It has

less impact on melt FeO_i and FeO_i/MgO than the nature of the protolith (Pg are more FeO_i rich than Og melts, Fig. 6). In the Og charge, orthopyroxene is present in negligible amounts. In the Pg charge, orthopyroxene is more abundant but other peritectic ferromagnesian phases (spinel, ilmenite) and residual biotite are also present and, so, the influence of orthopyroxene on melt composition is diluted. We speculate that the role of orthopyroxene would become more important at temperatures above 850 °C allowing biotite to be more extensively reacted and concentrations of ferromagnesian components to increase in the melt. To conclude, the major element characteristics and convergence of melt compositions generated from the two protoliths are best explained by the combined influence of the main melting reactions, i.e., reactions (1)–(3) act together to produce melts with a narrow range of major element compositions.

Trace element compositions of experimental melts

In contrast with the nearly homogeneous glass major element data, our melt trace element data show significant chemical variability, both at 800 and 850 °C. Assuming that analytical biases can be excluded, since contaminated glass spots have been filtered out, we explore below the possibility that the observed variability originates from heterogeneities in our starting materials. One possible type of heterogeneity is chemical, due to the variability of trace element concentrations in major phases of the protolith, mainly micas. Indeed, Rb (558–1034 ppm), Nb (87–228 ppm), Ta (3–10 ppm) and W (5–51 ppm) significantly vary in the Og biotite (Table 3). However, muscovite in Og, is less heterogeneous (74–88 ppm Li, 565–683 ppm Rb, 77–82 ppm Nb, 2–3 ppm Ta, 69–103 ppm W; Table 3). Chemical variability in melts (calculated as standard deviation/mean of analyses) reach 26% for Nb and 86% for Ta in the 800 °C charge (Table 5). These variations appear very elevated since Nb and Ta are nearly homogeneously distributed in muscovite (only 3% and 11% dispersions). Chemical dispersions for Li and Rb in melts (respectively, ~18% and ~13%) are also higher than the compositional variability in the starting muscovite (dispersions of ~9% and ~10%). Last, the dispersions for W in melts (~28% and ~22% at 800 °C and 850 °C) match the one in muscovite (~23%) but overall mica chemical heterogeneity seems unable to explain the glass variability for all trace elements.

A second type of heterogeneity in the natural protoliths is textural. In Og microcores, micas are segregated in few layers (Fig. 1). Melt pools are preferentially associated with muscovite reaction zones pre-localized in micaceous layers although mica-free zones also contain melt (see above). Melt films extend to lengths \geq ~1 mm and melt pools coexist at distances $>$ ~1 mm from each other (Fig. 4). Since, according to major elements (see above), melting is

kinetically controlled by dissolution of reactants and diffusive mass transport in the melt, chemical homogenisation of trace elements should take place diffusively. Diffusion distances in the melt (x) have been calculated for the different trace elements at 800 and 850 °C for a duration (t) of 14 days using the tracer diffusion (D) database of Zhang et al. (2010). Results ($x = (Dt)^{0.5}$) stress the large differences existing between trace elements. For Nb, Ta, Sn and W, very short diffusion distances were obtained, limited to 50 μm maximum (calculations performed with both dry and wet rhyolitic melt D). Maximum diffusion distances for Cs are 100 μm (calculations with dry and wet D) and they increase to $>$ 2 mm for Rb (dry and wet D) and $>$ 10 mm for Li (dry D). These results are broadly consistent with chemical dispersions in glasses being the lowest for Rb (~13 and ~10%) and Li (18 and 22%) and the highest for Ta (~86% and 41%) at 800 and 850 °C, while Nb (~26% and ~25%), Cs (~53% and ~24%) and W (~28% and ~22%, data in Table 5) are intermediate. Taken globally, these calculations support the possibility of incomplete diffusive homogenization between rare element-enriched and -depleted zones of melt films (respectively, near and away from reacted muscovites). However, for Li, the large diffusion distances calculated (in the same range as microcore sizes) are inconsistent with the observed dispersions and suggest that Li diffusive homogenization is perturbed by other processes, possibly Li incorporation in residual phases. We conclude that the trace element variability observed in experimental glasses is the consequence of combined chemical, mineralogical and textural heterogeneities inherited from the starting materials.

Sn is very low ($<$ 20 ppm) in all analyzed glasses and also in most glass–peritectic phase mixtures despite high concentrations in one protolith (~220 ppm Sn in the Og, Table 1). This uniform Sn depletion contrasts with the behavior of other elements, in particular Nb, Ta and W and is attributed to Sn loss by alloying with the Au capsule, an experimental difficulty particularly severe under reducing conditions as in this study (e.g., Linnen et al. 1995).

Behavior of rare elements during partial melting

Several models have been proposed to describe the behavior of trace element during partial melting (e.g., Shaw 1970; Hanson 1978; Harris and Inger 1992). Concentrations in melt relative to source rock can be calculated for variable F with a relatively small number of input parameters (source rock concentrations, mineral/melt partition coefficients). In most models, the restite remains mineralogically the same from the onset of melting until melt extraction, see Hertogen and Gijbels (1976) for an exception. In contrast, complex mineralogical changes take place in our experiments such as (i) disappearance of reactant phases (muscovite), (ii) appearance of peritectic

phases and (iii) changes in peritectic assemblage with varying temperature and melt fraction. These complexities have been circumvented by observing that the concentration of a given trace element relative to the source depends only on the bulk restite/melt partition coefficient (D) and F at the time of melt extraction (e.g., Hanson, 1978; Harris and Inger, 1992). This stresses the importance of D which determines whether a trace element is preferentially concentrated in the melt or sequestered in restite phases. Rare element enrichment in crustal melts requires $D < 1$.

D cannot be determined rigorously from our experiments because two types of information are lacking. First, peritectic phases have not been analyzed individually for trace elements although data on glass–peritectic phase mixtures are available. Second, the restite assemblage includes residual reactant phases (see “Introduction”) which should equilibrate with the formed melt but have not been characterized in experimental charges. We also assume that concentrations of trace elements in the melt are lower than the solubility of minerals with trace elements as ESCs (i.e., of cassiterite for Sn, see Linnen and Cuney 2005 for a summary of melt concentrations implied) so that no trace element mineral is part of the restite assemblage. Yet, two methods are presented below to constrain D for the rare elements of this study. Both are general in their principle but the Og 800 °C charge is selected for demonstration because (i) only a few trace element data are available for Pg charges, (ii) the 800 °C Og charge has the largest number of analyses of glass–peritectic phase mixtures and (iii) biotite is not significantly involved in melt production (at the difference of at 850 °C) and so muscovite dehydration-melting is the only reaction controlling the trace element distribution. In method 1, the restite is reduced to the peritectic assemblage (thus, $D \sim C_{\text{per}}/C_{\text{melt}}$ where C_{per} is the trace element concentration in the bulk peritectic assemblage). This assumes that residual phases do not equilibrate with the melt, which is reasonable given the short experimental durations and the slow element diffusivities in minerals. The peritectic assemblage is taken globally despite being in fact multiphase, i.e., including Bt_{per} , Spl_{per} , Ilm_{per} , Als_{per} and Kfs_{per} in reaction (3). For each trace element, C_{per} is calculated by subtracting the melt contribution from analyses of melt–peritectic phase mixtures, following the mass-balance equation:

$$C_{\text{mix}} = C_{\text{melt}} \cdot M_{\text{melt}} + C_{\text{per}} \cdot (1 - M_{\text{melt}}), \quad (4)$$

where C_{mix} and C_{melt} are as defined above and M_{melt} is the mass fraction of melt in the analyzed mixture. This latter is estimated from surfacic proportions of melt and peritectic phases determined on SEM images of mixtures zones taken prior and after LA-ICP-MS analysis (Fig. A3), using image

processing methods (see above and Table A7). For each element, C_{per} is calculated from the different C_{mix} analyses and the different local M_{melt} , C_{melt} being constant and equal to the median concentration in the 800 °C charge (Table 5). Results are detailed in Table A7. To more fully document the dispersion, $D (= C_{\text{per}}/C_{\text{melt}})$ are calculated from all C_{per} and C_{melt} analyzed in the Og 800 °C charge and the results are summarized in Table A8.

In method 2, a mass-balance approach anchored in the melting reaction (3) is implemented. Masses of trace elements released by the melting reaction are calculated and then distributed at equilibrium between the melt and the restite assemblage. For a given trace element, the mass released to the melt is equal to the sum of concentrations in reactants (Table 3; A3) each weighted proportionally to the stoichiometry of the melting reaction (Table 7). As a first step, the restite is reduced to the peritectic assemblage to allow comparison with method 1. However, the multiphase nature of the peritectic assemblage is fully recognized and the trace element concentration of each peritectic phase (c_{per}) is calculated from $C_{\text{melt}} \cdot K_{\text{d}}$, where K_{d} is the partition coefficient for the phase concerned. The calculations are performed element by element and iteratively. An initial trace element concentration is allocated to the melt and c_{per} calculated for all peritectic phases. The calculations are repeated with different C_{melt} until the mass of the trace element in melt and peritectic phases equals the mass released by the reaction. From this mass-balanced set of C_{melt} and c_{per} , C_{per} is calculated from the sum of c_{per} times mass for all peritectic phases in the assemblage. Finally, D is calculated from $C_{\text{per}}/C_{\text{melt}}$, where C_{melt} is the measured concentration (Table A9). Method 2 requires K_{d} for all trace elements in all phases of the peritectic assemblage. In this study, a set of partition coefficients derived from the Macusani Volcanics has been used (Table A9). Biotite/liquid K_{d} for Li, Rb, Nb, Cs, Ta (respectively, 3.40, 2.74, 9.69, 0.44, 2.07, Table A9) compare well with the results (1.7, 1.9, 4.85–9.21 0.3, 2.64–3.93) of Icenhower and London (1995) and Stepanov and Hermann (2013). K -feldspar/liquid K_{d} for Li, Rb and Cs (respectively, 0.12, 0.95 and 0.026, Table A9) are within literature ranges (0.05–0.22, 0.68–1.03 and 0.03–0.13, Icenhower and London 1996; Acosta-Vigil et al. 2010; Hulsbosch et al. 2014; Villarros and Pichavant 2019). For Al silicate (Als_{per}), the K_{d} (andalusite/melt data) are all in the 0.01 to 0.001 range (Table A9). Hercynitic spinel (Spl_{per}), the only phase for which the K_{d} were estimated (Table A9), is assumed to be the W-bearing phase required by the C_{mix} data (see above).

C_{per} and D calculated with both methods are plotted in Figs. 8 and 9, respectively. The dispersion in C_{per} with method 1 is the result of several factors including differences in peritectic modal compositions (see above, Fig. A3 and Table A7), mineralogical variations in the third

direction (calculations of M_{melt} only use 2D images) and spatial variability in C_{melt} . As a consequence, D calculated with method 1 are dispersed and, for some elements, significantly different D are obtained with method 1 when compared to method 2 (Fig. 9). However, the important point is that the two methods yield consistent partitioning trends for all elements.

Li, Cs. These two elements show a similar behavior. Both the C_{mix} data (see above) and the C_{per} and D calculations consistently demonstrate strong preferential partitioning toward the melt. Bt_{per} is the only major host for Li ($K_{\text{dLi}} > 1$, Table A9) and for Cs in the peritectic assemblage, being only present in small proportions (Table 7). Therefore, for both elements, D stays < 1 (Fig. 9). However, D_{Li} higher than in this study are expected for protoliths and muscovite dehydration-melting reactions leading to higher Bt_{per} proportions than found in the Og peritectic assemblages. Therefore, for melts generated by muscovite dehydration-melting, Li concentrations are inversely correlated with proportions of biotite in the peritectic assemblage. In cases where the melt has time to equilibrate with residual Li-bearing phases such as biotite before being extracted, an increase in D_{Li} relative to values in this study is also anticipated, which would reduce the melt Li content.

Rb. Both the C_{mix} data and the C_{per} calculations show that Rb is not very much fractionated between melt and peritectic assemblage. D_{Rb} is lower than 1 (Fig. 9) with the two calculation methods indicating preference for the melt. With K_{dRb} for Kfs_{per} slightly < 1 and K_{dRb} for other peritectic phases $< < 1$, Bt_{per} ($K_{\text{dRb}} > 2.5$, Table A9) is again not abundant enough in the 800 °C Og charge to increase C_{per} and so D_{Rb} is pulled to values < 1 . Elimination of K-feldspar from the peritectic assemblage (for example, if melting takes place under fluid present, Harris and Inger 1992) would further decrease D_{Rb} and increase the Rb concentration of the melt. However, melt fractions generally differ between fluid present and fluid absent. The stronger Rb enrichment in melt found under fluid absent (Harris and Inger 1992) is most probably the result of F being much smaller in the former (0.11) than in the latter case (0.41) since D_{Rb} should be higher. This illustrates the interplay between the two variables, F and D , in the control of trace element enrichment.

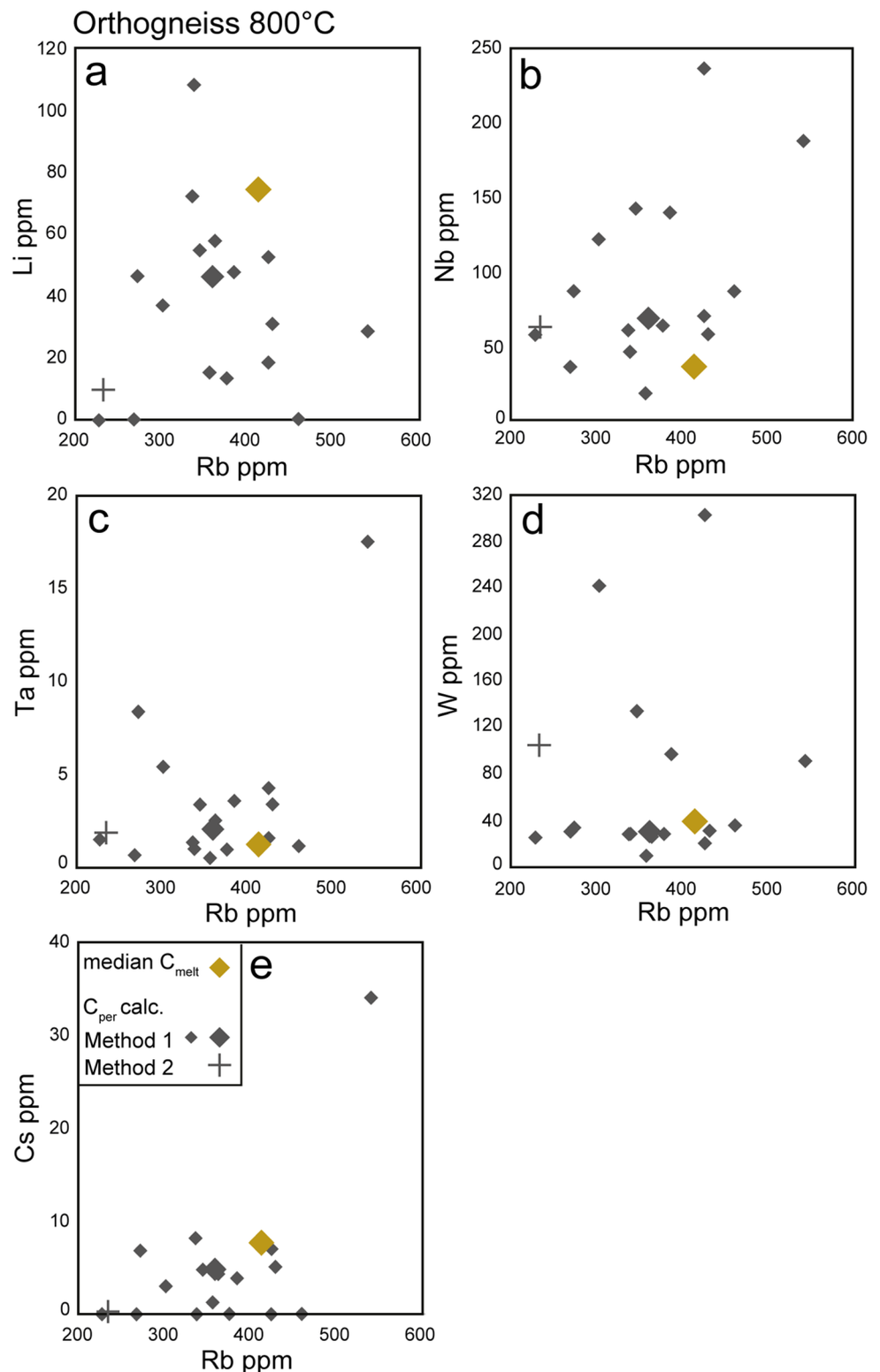
Nb, Ta. For these two elements, consistent D values are obtained with the two methods, confirming the C_{mix} data and indicating preferential partitioning toward the peritectic assemblage. The measured and calculated C_{melt} are also very close to each other (Table A9) but we point out that the calculated C_{melt} sensitively depend on K_{dNb} and $K_{\text{dT a}}$ for Spl_{per} . Fluid-absent dehydration-melting of muscovite results in Nb and Ta mainly sequestered in peritectic phases. However, our data do not allow to conclude on melt/peritectic Nb/Ta partitioning, a consequence of the dispersion in C_{melt} and C_{per} .

The median Nb/Ta (higher in C_{per} than in C_{melt} , Tables 5, A7, A9; Fig. 10b) suggest that Nb is slightly more preferentially partitioned than Ta in the peritectic assemblage but the C_{mix} data (see above) and D calculations (Tables A8; A9) imply the opposite. In any case, the melt/peritectic Nb/Ta fractionation is small (Stepanov and Hermann 2013; Stepanov et al. 2014). Melts produced by dehydration-melting of muscovite in Og charges have Nb/Ta (medians ~ 27 – 29 , Table 5) significantly higher than the Og source rock (Nb/Ta = 12, Table 1) and mainly reflect the composition of the starting muscovite (Nb/Ta ~ 32 – 39 , Fig. 10b). Despite the presence of Bt_{per} (Nb/Ta ~ 140 from Table A9) in the 800 °C charge, the melt Nb/Ta is not pulled to values much lower than muscovite because the proportion of Bt_{per} is small and counterbalanced by other low Nb/Ta peritectic phases. However, as emphasized above for Li, a Bt_{per} proportion higher than observed in this study would presumably lead to a Nb/Ta restite–melt fractionation (Stepanov and Hermann 2013) stronger than observed in Og charges. Nb and Ta, thus, provide further illustration of the importance of peritectic phase assemblages and proportions for element partitioning during crustal melting.

W. From the C_{mix} data, this element is commonly but not systematically enriched in glass–peritectic phase mixtures (see above) and the D_{W} results are dispersed. With method 1, D_{W} ranges from 0.64 to ~ 3 (median slightly < 1 , Table A8). With method 2, D_{W} (2.70) is in the upper range of results with method 1 (Table A9; Fig. 9). Although caution is needed with method 2 which sensitively depends on K_{dW} for Spl_{per} , calculated and measured C_{melt} are in close agreement from each other (Table A9). In other words, changing K_{dW} for Spl_{per} would lead to calculated C_{melt} differing from measured. It is also reminded that, at 850 °C, the C_{mix} data consistently indicate W enrichment in mixtures (see above). Therefore, the high D_{W} results (upper part of the range with method 1 and method 2) are preferred and values < 1 with method 1 are interpreted to represent Spl_{per} -poor peritectic mixtures. This implies that W is partitioned in the peritectic assemblage rather than in the melt. We note that K_{dW} for biotite is < 1 (Table A9) unlike other elements (Li, Rb, Nb, Ta) and so large changes in D_{W} are not expected in case of Bt_{per} -rich peritectic assemblages. This implies that biotite is not a key W sequestering phase in high grade residual source rocks. In contrast, the elevated K_{dW} for muscovite (~ 2 , Raimbault and Burnol 1998; Michaud and Pichavant 2020) makes this phase critical in the supply of W to crustal melts.

Sn. Because of Sn loss to the capsule and underestimation of C_{melt} and C_{mix} , method 1 cannot be implemented. Results with method 2 (calculated $C_{\text{melt}} = 50.8$ ppm, $D = 0.36$, Table A9) depend directly on K_{dSn} for Spl_{per} which cannot be constrained from C_{melt} (as done for W) and so the partitioning behavior of Sn remains unconstrained.

Fig. 8 Trace element concentrations in peritectic assemblages (C_{per}) calculated for the Og charge at 800 °C. Results with the methods of calculation detailed in text are compared (method 1: black diamonds; method 2: cross). For method 1, the small symbols represent individual calculations and the large symbol the median value (Table A7). Results with method 2 are detailed in Table A9. Median trace element concentrations in melt (C_{melt} ; Table 5) are shown for comparison



Rare element distributions during biotite dehydration-melting are not as well constrained as during muscovite dehydration-melting. However, method 2 was tentatively applied to the 850 °C Pg and reaction (2) assuming K_{dLi} and

K_{dCs} for orthopyroxene = 0. It yields a calculated C_{melt} of 164 ppm for Li (vs. 183 ppm, Table 5) and of 27 ppm for Cs (vs. 19 ppm, Table 5), consistent with biotite (163–219 ppm Li, 29–31 ppm Cs, Table 3) being the main reactant in (2).

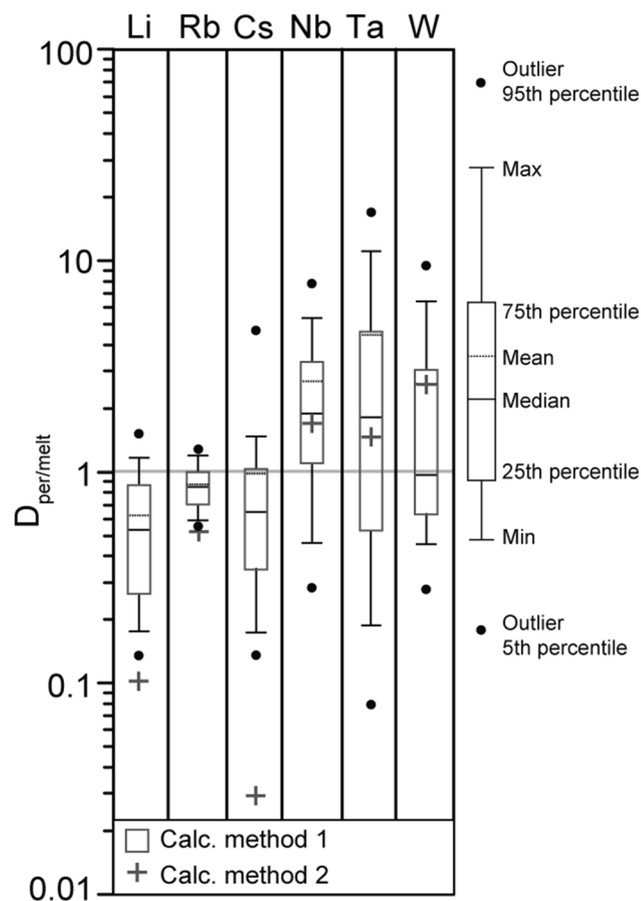


Fig. 9 Bulk trace element partition coefficients (D) between melt and peritectic assemblages for Li, Rb, Cs, Nb, Ta, W. Results using method 1 are shown as boxplots (Table A8) and using method 2 as crosses (Table A9). D values > 1 indicate preferential partitioning toward the peritectic assemblage and $D < 1$ toward the melt. See text for additional details

Therefore, Li concentrations in melts generated by biotite dehydration-melting are positively correlated with proportions of reactant biotite and its Li concentrations. Both the calculated D_{Li} and D_{Cs} are very low (< 0.1), consistent with the inferred absence of Li- and Cs-bearing phase in the peritectic assemblage. However, as noted for muscovite dehydration-melting, chemical equilibration of the melt with residual biotite (if present) would increase the D values.

Implications for natural rare element-rich melts

Comparison between experimental melts and Variscan granites

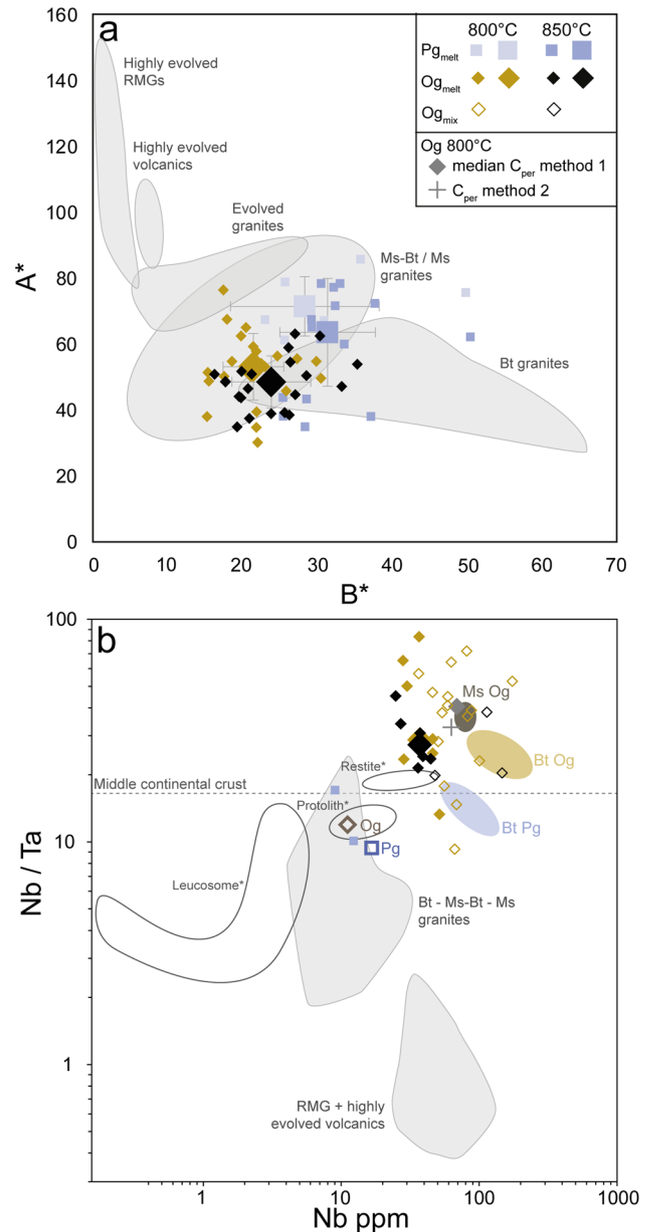
Major element chemistries of experimental melts are compared with whole-rock compositions of Variscan

granites and volcanic equivalents with examples taken from Cornwall (Simons et al. 2016, 2017; Manning and Hill 1990), Portugal (Charoy and Noronha 1996; Neiva 2002; Michaud 2019; Michaud and Pichavant 2020), French Massif Central (Raimbault et al. 1995; Raimbault and Burnol 1998; Villaros and Pichavant 2019), Armorican Massif (Ballouard et al. 2015), Spain (Chicharro et al. 2016), Erzgebirge (Breiter et al. 2005) and from the Macusani Volcanics (Pichavant et al. 1987). Variscan granites include peraluminous biotite, biotite-muscovite, muscovite and rare metal granites. They exhibit various degrees of fractionation marked, on major elements, by an increase of A^* at decreasing B^* (Fig. 10a). For the comparison with melts to be meaningful, granite whole rocks should be representative of their parental liquid compositions. Although this might not be the case for all examples in Fig. 10, peraluminous biotite, biotite-muscovite, muscovite leucogranites and RMGs are generally considered to be emplaced as almost pure liquids (Raimbault et al. 1995; Scaillet et al. 1995; Patiño Douce and Harris 1998; Villaros and Pichavant 2019). All experimental melts from this study plot in the field of peraluminous muscovite-biotite/muscovite granites (Fig. 10a). Again, no major difference emerges with the nature of the protolith implying that leucogranites with similar major element chemistries can be generated by melting of either an orthogneissic or a paragneissic source rock (see the discussion on major element compositions). The Pg melt compositions do not extend into the high B^* domain characteristic of biotite granites (Fig. 10a). This is the consequence of biotite being part of the residual phase assemblage in the 850 °C Pg which buffers the concentration of ferromagnesian components in the melt (see above). No partial melt from this study approaches the most strongly peraluminous and leucocratic major element compositions characteristic of RMGs and volcanic equivalents (field with highest A^* and lowest B^* in Fig. 10a).

In comparison with the reference middle continental crust, most Variscan granites are rare element enriched. The evolution from peraluminous biotite, biotite-muscovite, muscovite leucogranites to RMGs and volcanic equivalents is marked by increasing Nb, Ta and Li at increasing Rb (Fig. 10b, 11). In some peraluminous granites, W reaches high concentration levels, in the RMG range (Fig. 11c). Peraluminous granites and RMGs plot below the reference Nb/Ta of the middle continental crust, the chemical evolution from the least to the most enriched compositions being marked by a strong decrease of Nb/Ta with increasing Nb (Fig. 10b, Stepanov et al. 2014). For some elements (Li, Rb, Ta), concentrations in experimental glasses (Pg and Og) and in the least evolved peraluminous granites overlap. However, the most fractionated rocks are way more enriched than the experimental

Fig. 10 Comparison between experimental melts from this study (Tables 4, 5) and compositions of Variscan granites and volcanic equivalents (Cornwall: Simons et al. 2016, 2017; Manning and Hill 1990; Portugal: Charoy and Noronha 1996; Neiva 2002; Michaud 2019; Michaud and Pichavant 2020; French Massif Central: Raimbault et al. 1995; Raimbault and Burnol 1998; Villaros and Pichavant, 2019; Armorican Massif: Ballouard et al. 2015; Spain: Chicharro et al. 2016; Erzgebirge: Breiter et al. 2005; Macusani Volcanics: Pichavant et al. 1987). **a** Major element compositions plotted in the A^* - B^* diagram of Debon and Lefort (1983), see Tables 1 and 4. **b** Nb/Ta vs. Nb content. In both diagrams, the C_{melt} data (either for the Pg, Pg_{melt} or for the Og, Og_{melt}) are plotted using the same color coding as in Fig. 7, the small symbols representing individual analyses and the large symbols median values (Table 5). In (b), the C_{mix} individual analyses for the orthogneiss (Og_{mix}) at 800 and 850 °C (Table 6) and the C_{per} data for the orthogneiss at 800 °C (Og 800 °C) are also plotted for comparison (method 1: median from Table A7 and method 2 from Table A9). The large open blue square and black diamond, respectively, labeled Pg and Og represent the whole-rock compositions of the two protoliths. Compositions of starting muscovite in the orthogneiss (Ms Og) and of starting biotite in the orthogneiss (Bt Og) and paragneiss (Bt Pg) are shown. The horizontal dashed line gives the mean Nb/Ta composition of the middle continental crust (Rudnick and Gao 2003). Fields for protoliths, leucosomes and restites from the Ivrea zone, Italy, all labeled with an asterisk, are indicated for reference (Data from Bea and Montero 1999 see also Stepanov and Hermann 2013)

glasses (Fig. 11). Differences are large and the most enriched experimental glasses have Li and Ta concentrations about ~ 1 order of magnitude lower than RMGs. The Nb contents of the least enriched (Pg) glasses are similar to peraluminous granites whereas the most enriched (Og) are in the RMG range (Fig. 11a). W is the other element whose experimental concentrations (particularly in Og) are comparable to concentrations in RMGs and volcanic equivalents (Fig. 11c). The Nb/Ta of Pg melts plots with the least evolved peraluminous granites, similar to leucosomes in granulite facies metapelites (Bea and Montero, 1999) and slightly below the middle continental crust and restites from biotite dehydration-melting (Fig. 10b). In comparison, the Nb/Ta of Og melts are higher than the range of natural granites and also than restites (Fig. 10b). As for the major elements, the trace element characteristics of the melts are the direct result of melting mechanisms in the experiments. The high Nb, W, Nb/Ta of Og melts generated by muscovite dehydration-melting reflect the major role of muscovite in the supply of these elements. The small abundance of peritectic biotite in Og charges (a consequence of the Og felsic bulk composition) limits biotite/melt Nb/Ta fractionation (i.e., preferential sequestration of Nb than Ta in biotite, Stepanov and Hermann 2013) and allows the melt Nb concentration and Nb/Ta to remain high, not so different from in the starting muscovite (Figs. 10b, 11). In contrast, the Nb/Ta of Pg melts at 850 °C is mainly controlled by biotite dehydration-melting, both melt and biotite having similar Nb/Ta ratios (Fig. 10b). As biotite persists as a residual phase,



Nb remains preferentially sequestered in biotite and melts are not Nb-enriched, in good agreement with Stepanov and Hermann (2013). The Nb depletion in leucosomes when compared to amphibolite grade metasediments ("protolith" in Fig. 10b) is also consistent with biotite being involved either as a peritectic or a residual phase. Similarly, the higher Li concentrations in Pg than in Og melts reflect the respective proportions of reactant biotite in (2) and (3). Overall, and while noting that caution is required with the use of whole-rock data as proxies of melt concentrations (see Michaud and Pichavant 2020 for illustration), significant differences exist between the rare element patterns of experimental melts and Variscan granites.

Influence of rare element pre-concentration

Differences between experimental and natural compositions (Figs. 10, 11) can be interpreted in various ways, one possible explanation revolving around the nature of the protoliths used in the experiments. On the one hand, our Og protolith is somewhat atypical as it leads to peritectic assemblages relatively poor in Bt_{per} which impacts the Li, Cs, Nb, Ta distributions and Nb/Ta fractionation (see above). Nevertheless, the important point is that mechanisms of rare element distribution during muscovite dehydration-melting have been clarified from the Og and can now be applied to other protoliths such as metapelites. On the other hand, it has been commonly proposed that specific rare element pre-enriched protoliths are required for the production of evolved rare element-enriched granitic melts (Romer and Kroner, 2015; Ballouard et al. 2018; Roda-Robles et al. 2018) and it is worth recalling that the Og is representative of a such a pre-concentrated source. Yet, glasses and glass-peritectic phase mixtures generated from the Og are not uniformly enriched in rare elements. For W, the high concentrations in experimental melts (medians ~ 30–40 ppm, Table 5) and peritectic mixtures (up to ~ 200 ppm, Table 6) match the W enrichment in the source rock (~ 17 ppm, Table 1; Fig. 11c) but the high Nb and Ta concentrations have no equivalence in the bulk source (Fig. 11a, b) and the Li (Fig. 11d) and Cs melt concentrations are not very high despite their enrichments in the Og. We also point out that melts with the highest Li (168–198 ppm in the 850 °C Pg charge, Table 5; Fig. 11d) derive from the least Li-rich protolith. Thus, source pre-concentration is not the only process controlling the trace element contents of anatectic melts.

Mobilization of rare elements depends on proportions of reactants in the melting reaction and on their rare element concentrations. For a given melting reaction and P–T conditions (hence fixed reactant proportions), the amount of mobilized rare elements increases with the rare element concentration of reactants. Therefore, a source rock with “enriched” reactant phases will supply a higher amount of rare elements than a barren protolith. However, melting reactions and proportions of reactants are specific of the protolith, sensitively depending on its bulk chemistry and mineralogy as illustrated in Table 7. In addition, mobilization is only one step in the overall process and so a high amount of mobilized rare elements does not necessarily result in their complete transfer to the melt. This is best illustrated by Og experiments which show that, upon muscovite dehydration-melting, Nb and Ta are preferentially sequestered in the peritectic assemblage (Fig. 11a, b). In contrast, Li and Cs are strongly partitioned toward the melt. For those elements, melt–peritectic partitioning has a lesser role and source pre-enrichment is potentially more important than for Nb and Ta. We conclude that rare element pre-concentration has some influence on

enrichment levels attainable in the melt. However, this influence is highly selective between rare elements and we caution against source pre-concentration being viewed as the only factor controlling the generation of “enriched” crustal melts. The type of melting reaction, proportions of reactants, their individual rare element concentrations and restite–melt partitioning are equally important.

Implications for rare element-rich melt generation

Another possible explanation for the geochemical differences between experimental melts and natural granites (Figs. 10, 11) concerns melting mechanisms. An anatectic origin is generally considered for the least evolved crustal peraluminous magmas but also, in the frame of the crystallization-differentiation model, for magmas parental to RMGs and LCT pegmatites (e.g., Raimbault et al. 1995; Breaks et al. 2005; Černý et al. 2005; Roda-Robles et al. 2018; Villaros and Pichavant, 2019). Our results confirm that mica dehydration-melting (i.e., reactions (3) and (2)) can produce peraluminous crustal melts similar to Variscan muscovite granites (Fig. 10, Patiño Douce and Harris 1998). However, these melting mechanisms apparently fail to generate the most peraluminous, leucocratic and rare element-enriched compositions. In the following we discuss (1) fluid-present melting and (2) residual source melting.

According to the first model, devolatilization of the lower crust generates carbonic fluids rich in F and Cl which are also transporting agents for Li, Cs and Sn and explain their large-scale depletions in granulites (Cuney and Barbey 2014 and references therein). Influx of deep fluids promotes partial melting at intermediate crustal levels leading to the generation of F- and rare metal-rich granitic melts (Cuney and Barbey 2014). Despite some debatable points such as the fluid control of element fractionation in the lower crust (see Stepanov and Hermann 2013 for an alternative interpretation), this fluid-present model accounts for many geological and geochemical features of rare metal magmatism in orogenic belts such as the Variscan. However, unfortunately, experimental information is still almost totally lacking on fluid-present melting, defined here as melting of crustal protoliths in presence of a fluid phase of complex and variable compositions (Montel et al. 1986; Johannes and Holtz 1996; Villaros et al. 2018b). Experimental results from this study contribute very little on this mechanism since the exploratory fluid-present experiments mentioned earlier had high melt fractions inconsistent with degrees of melting for peraluminous granites and the fluid-present reaction (1) is only important near the solidus. To discuss the behavior of major and trace elements during fluid-present melting, the systematic determination of melting reactions, phase assemblages, melt compositions and fluid/melt partition coefficients as a

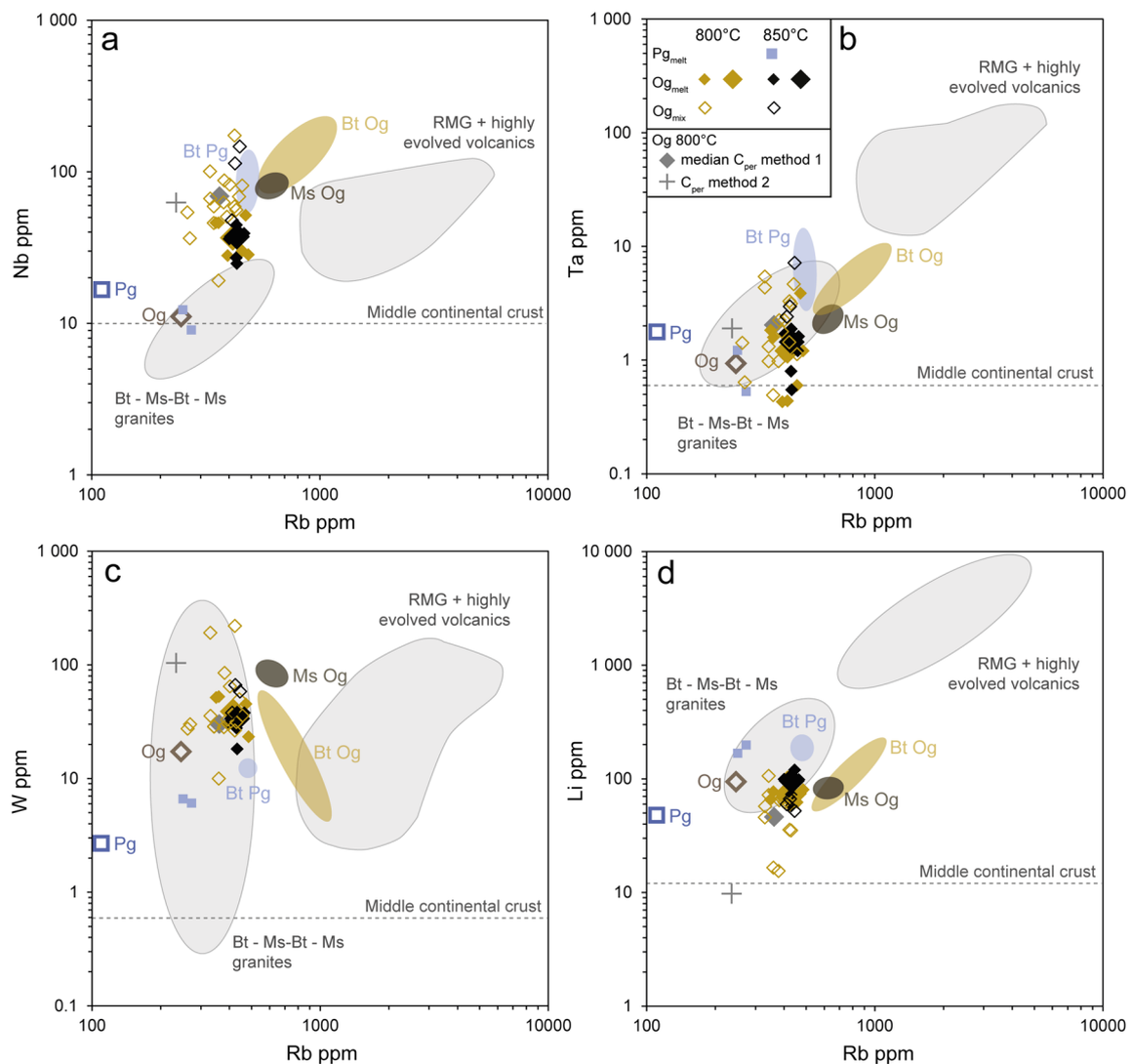


Fig. 11 Comparison between experimental melt from this study (Table 5) and trace element compositions of Variscan granites and volcanic equivalents (same natural examples as in Fig. 10). **a** Nb, **b** Ta, **c** W, **d** Li concentrations plotted as a function of Rb. In all diagrams, the C_{melt} data (either for the Pg, Pg_{melt} or for the Og, Og_{melt}) are plotted using the same color coding as in Fig. 6, the small symbols representing individual analyses and the large symbols median values (Table 5). In (**b**), the C_{mix} individual analyses for the orthogneiss (Og_{mix}) at 800 and 850 °C (Table 6) and the C_{per} data for the

orthogneiss at 800 °C ($Og_{800\text{ °C}}$) are also plotted for comparison (method 1: median from Table A7 and method 2 from Table A9). The large open blue square and black diamond, respectively, labeled Pg and Og represent the whole-rock compositions of the two protoliths. Compositions of starting muscovite in the orthogneiss (Ms Og) and of starting biotite in the orthogneiss (Bt Og) and paragneiss (Bt Pg) are shown. The horizontal dashed lines give the mean compositions of the middle continental crust (Rudnick and Gao 2003)

function of pressure, temperature and fluid composition is needed.

In the second model, illustrated below using the Variscan belt as an example, the melting scenario involves two successive melting events (e.g., Wolf et al. 2018). First, conditions of muscovite dehydration-melting are reached in the continental crust and muscovite-bearing lithologies undergo partial melting. P–T paths show a generalized melting event at ~330 Ma in the Variscan belt (Faure et al. 2005, Gébélín et al. 2007, Le Carlier de Veslud et al. 2013), under

conditions compatible with the generation of biotite–muscovite/muscovite peraluminous granites by muscovite dehydration-melting (FMC: Melleton et al. 2009, Faure et al. 2005, Lardeaux 2014; Cornwall: Simons et al. 2016). Following reaction (3), rare elements are released and distribute between the generated melt and the restitic assemblage. Depending on mechanisms of melt extraction, variable levels of enrichment in ascending magmas are generated for rare elements mobilized during muscovite dehydration-melting such as W. This provides a general explanation for the spatial

association between biotite–muscovite/muscovite peraluminous granites and hydrothermal W-deposits, both in the Variscan belt (e.g., Cornwall: Simons et al. 2016; FMC: Harlaux et al. 2018) and beyond (e.g., China: Yuan et al. 2019; Rwanda: Hulsbosch 2019). Second, conditions of biotite dehydration-melting are attained in some specific parts of the belt most probably in response to local thermal anomalies (e.g., Velay: Barbey et al. 2015, Villaros et al. 2018a; Cornwall: Simons et al. 2016). Restitic biotite-bearing lithologies (made of residual plus peritectic phases from the first melting event), undergo partial melting. Our results show that the peritectic assemblage left after muscovite dehydration-melting is enriched in Nb, Ta, W compared to the initial protolith, the behavior of Sn being still uncertain (see above, Wolf et al. 2018). Enrichment of Li in the peritectic assemblage, although not observed in this study, would go along with the proportion of peritectic or residual biotite and is to be expected for protoliths derived from mica-rich metapelites. Such a residual, rare element-enriched assemblage makes an appealing source for the generation of RMG-like melts. Biotite dehydration-melting would, thus, result in the production of Li-, Nb-, Ta- and W-rich melts with low Nb/Ta although detailed information on melting reactions, phase assemblages and melt compositions are still necessary. In particular, it remains to be demonstrated that the residual source model can lead to chemically specific melts, both for major (strongly peraluminous and leucocratic) and rare elements (strongly enriched in Li, Ta, Fig. 11).

Our experimental results and the evaluation of alternatives to mica dehydration-melting emphasize difficulties with the genesis of RMG-like melts by purely anatectic processes. However, RMGs might be produced by a combination of petrogenetic processes rather than by partial melting alone. For example, crystallization differentiation might act together with mica dehydration-melting to produce the distinctive RMG geochemical signatures (e.g., Pichavant et al. 1988b; Raimbault et al. 1995; Roda-Robles et al. 2018; Michaud and Pichavant 2020). Evaluation of this scenario requires combined melting-fractionation models incorporating the type of information presented in this study for partial melting processes.

Conclusions

In this study, we have emphasized the role of melting mechanisms and experimentally documented the behavior of representative rare elements upon micas dehydration-melting in the continental crust. Despite the contrasted nature and mineral modal abundances of the two protoliths, and the different melt distributions, melt major element compositions are nearly homogeneous, a result of the combined influence of the main melting reactions. In contrast, the significant

variability of melt trace element compositions originates from heterogeneities in the starting materials (i.e., mineral compositions and distribution).

The amounts and the behavior of rare elements released upon melting is controlled by the stoichiometry of the reaction, the nature, proportions and compositions of reactant phases and the partition coefficients between melt, residual and peritectic phases. Rubidium, Nb, Ta and W are enriched and Li and Cs depleted in $O_{\Sigma \text{melts}}$ compared to source rock while $P_{\Sigma \text{melts}}$ are strongly enriched in Li and also in Rb, Cs, and W. Upon muscovite dehydration-melting, Li and Cs strongly and Rb slightly partition into the melt, whereas Nb, Ta and W partition to the peritectic assemblage. Upon partial biotite dehydration-melting, limited data indicate a partitioning of Li and Cs into the melt in contrast to Nb and Ta. The modal proportion of biotite, either peritectic or residual, strongly affects the behavior of Li, Rb, Nb, Ta but not of W, and the Nb/Ta.

Melt major element compositions are similar to those of muscovite of the Variscan belt of Western Europe but strongly differ from RMGs and volcanic equivalents. In addition, trace element patterns are also contrasted between experimental melts and natural compositions. These mismatches might partly reflect the specificities of the starting materials. Though, the persisting issues with genesis of RMG-like melts through partial melting alone suggest that petrogenetic processes such as crystallization differentiation might be combined to micas dehydration-melting. Besides, fluid-present and multistage melting remain to be tested experimentally as alternatives.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s00410-021-01855-9>.

Acknowledgements This study was initiated as part of the doctoral thesis of JM supported by the Labex VOLTAIRE (ANR-10-LABX-100-01), the ERAMIN project NewOres and the ANR project VARPEG (ANR-15-CE01-0001). The authors acknowledge Dr. I. Di Carlo for assistance with analyses at the microprobe. The manuscript significantly benefited from the relevant recommendations and comments of the Editor in Chief Pr. Dr. Othmar Müntener, Pr. Dr. Robert Linnen and an anonymous reviewer.

References

- Acosta-Vigil A, Buick I, Hermann J, Cesare B, Rubatto D, London D, Morgan GB (2010) Mechanisms of crustal anatexis: a geochemical study of partially melted metapelitic enclaves and host dacite, SE Spain. *J Petrol* 51:785–821
- Albarède F (1995) Introduction to geochemical modeling. Cambridge University Press, Cambridge, p 543
- Ballouard C, Boulvais P, Poujol M, Gapais D, Yamato P, Tartèse R, Cuney M (2015) Tectonic record, magmatic history and hydrothermal alteration in the Hercynian Guérande leucogranite, Armorican Massif, France. *Lithos* 220–223:1–22

- Ballouard C, Poujol M, Zeh A (2018) Multiple crust reworking in the French Armorican Variscan belt: implication for the genesis of uranium-fertile leucogranites. *Int J Earth Sci* 107:2317–2336
- Barbey P, Villaros A, Marignac C, Montel J-M (2015) Multiphase melting, magma emplacement and P–T-time path in late-collisional context: the Velay example (Massif Central, France). *Bull Soc Géol France* 186(2–3):93–116
- Bea F, Montero P (1999) Behavior of accessory phases and redistribution of Zr, REE, Y, Th, and U during metamorphism and partial melting of metapelites in the lower crust: an example from the Kinzigite Formation of Ivrea-Verbano, NW Italy. *Geochim Cosmochim Acta* 63:1133–1153
- Bea F, Pereira MD, Stroh A (1994) Mineral/leucosome trace-element partitioning in a peraluminous migmatite (a laser ablation-ICP-MS study). *Chem Geol* 117:291–312
- Belkamsi M, Cuney M, Pollard PJ, Bastoul A (2000) Chemistry of the Ta-Nb-Sn-W oxide minerals from the Yichun rare metal granite (SE China): genetic implications and comparison with Moroccan and French Hercynian examples. *Mineral Mag* 64:207–523
- Breaks F W, Selway J B, Tindle A C (2005) Fertile peraluminous granites and related rare-element pegmatites, Superior Province of Ontario. In: Linnen RL, Sampson IM (eds) Rare-element geochemistry and mineral deposits. *Geol. Soc. Can. Short Course Notes*, St. Catharines, 17:87–125
- Brearley FW (1987) An experimental and kinetic study of the breakdown of aluminous biotite at 800 °C: reaction microstructures and mineral chemistry. *Bull Miner* 110:513–532
- Brearley AJ, Rubie DC (1990) Effects of H₂O on the Disequilibrium breakdown of muscovite+quartz. *J Petrol* 31(4):925–956
- Breiter K, Müller A, Leichmann J, Gabašova A (2005) Textural and chemical evolution of a fractionated granitic system: the Podlesí stock, Czech Republic. *Lithos* 80:323–345
- Brown GC, Fyfe WS (1970) The production of granitic melts during ultrametamorphism. *Contrib Mineral Petrol* 28:310–318
- Burnham CW (1979) Magma and hydrothermal fluids. In: Barnes HL (ed) *Geochemistry of hydrothermal ore deposits*, 2nd edn. Wiley Interscience, New York, pp 71–136
- Cameron EN, Jahns RH, McNair AH, Page LR (1949) Internal structure of granitic pegmatites. *Econ Geol Monograph* 2:155p
- Carignan J, Hild P, Mevelle G, Morel J, Yeghicheyan D (2001) Routine analyses of trace elements in geological samples using flow injection and low pressure on-line liquid chromatography coupled to ICP-MS: a study of geochemical reference materials BR, DR-N, UB-N, AN-G and GH. *Geostand Newsl* 25:187–198
- Carlier Le, de Veslud C, Alexandre P, Ruffet G, Cuney M, Cheilletz A (2013) A two-stage exhumation in Western French Massif Central: new geochronological evidences of syn-collisional extension. *Lithos* 175–176:1–15. <https://doi.org/10.1016/j.lithos.2013.04.013>
- Černý P (1991) Rare-element granitic pegmatites. Part 1: anatomy and internal evolution of pegmatite deposits. Part 2: regional to global environments and petrogenesis. *Geosci Canada* 18:49–81
- Černý P (1992) Geochemical and petrogenetic features of mineralization in rare-element granitic pegmatites in the light of current research. *Appl Geochem* 7:393–416
- Černý P, Ercit TS (2005) The classification of granitic pegmatites revisited. *Can Mineral* 43:2005–2026
- Černý P, Masau M, Goad BE, Ferreira K (2005) The Greer Lake leucogranite, Manitoba, and the origin of lepidolite-subtype granitic pegmatites. *Lithos* 80:305–321
- Cesare B (2000) Incongruent melting of biotite to spinel in a quartz-free restite at El Joyazo (SE Spain): Textures and reaction characterization. *Contrib Miner Petrol* 139:273–284
- Chappell BW, White AJR, Wyborn D (1987) The importance of residual source material (restite) in granite petrogenesis. *J Petrol* 28:1111–1138
- Charoy B, Noronha F (1996) Multistage growth of a rare-element, volatile-rich microgranite at Argemela (Portugal). *J Petrol* 37:73–94
- Chicharro E, Boiron M-C, López-García JÁ, Barfod DN, Villaseca C (2016) Origin, ore forming fluid evolution and timing of the Logrosán Sn–(W) ore deposits (Central Iberian Zone, Spain). *Ore Geol Rev* 72:896–913
- Christiansen EH, Sheridan MF, Burt DM (1986) The geology and geochemistry of Cenozoic topaz rhyolite from the western United States. *Geol Soc Am Spec Paper* 205:1–82
- Clemens JD, Stevens G (2016) Melt segregation and magma interactions during crustal melting: breaking out of the matrix. *Earth Sci Rev* 160:333–349
- Clemens JD, Vielzeuf D (1987) Constrains on melting and magma production in the crust. *Earth Planet Sci Lett* 86:287–306. [https://doi.org/10.1016/0012-821X\(87\)90227-5](https://doi.org/10.1016/0012-821X(87)90227-5)
- Connolly JAD, Holness MB, Rushmer T, Rubie DC (1997) Reaction-induced micro-cracking: an experimental investigation of a mechanism for enhancing anatexis melt extraction. *Geology* 25:591–594. [https://doi.org/10.1130/0091-7613\(1997\)025%3c0591:RIMAEI%3e2.3.CO;2](https://doi.org/10.1130/0091-7613(1997)025%3c0591:RIMAEI%3e2.3.CO;2)
- Crevola G (1987) Les orthomicaschistes, produits de la déformation cisailante ductile symmétamorphe de granites avec transfert de matière: principaux caractères et exemples dans la chaîne Varisque du Sud de la France. *Geodin Acta* 1:207–221
- Cuney M, Barbey P (2014) Uranium, rare metals, and granulite-facies metamorphism. *Geosci Front* 5:729–745
- Cuney M, Marignac C, Weisbrod A (1992) The Beauvoir topaz-lepidolite albite granite (Massif Central, France); the disseminated magmatic Sn-Li-Ta-Nb-Be mineralization. *Econ Geol* 87:1766–1794
- Debon F, Le Fort P (1983) A chemical-mineralogical classification of common plutonic rocks and associations. *Trans Roy Soc Edinburgh* 73:135–149
- Deveaud S, Gumiaux C, Gloaguen E, Branquet Y (2013) Spatial statistical analysis applied to rare-element LCT-type pegmatite fields: an original approach to constrain faults-pegmatites-granites relationships. *J Geosci* 58:163–182. <https://doi.org/10.3190/jgeosci.141>
- Devineau K, Pichavant M, Villiéras F (2005) Melting kinetics of granitic powder aggregates at 1175°C, 1 atm. *Eur J Mineral* 17:387–398
- Falster AU, Simmons Wm B, Webber KL (1997) The origin of evolved LCT-type granitic pegmatites in the Hoskin Lake granite-pegmatite field, Florence Co., Wisconsin. IAVCEI General Assembly, Jan. 1997, Puerto Vallarta, Mexico, Programme, pp 118
- Faure M, BéMézème E, Duguet M, Cartier C, Talbot J-Y (2005) Paleozoic tectonic evolution of medio-europa from the example of the French massif central and massif armoricain. *J Virt Explor Electron Edit* 19(5):1–26
- Faure M, Lardeaux J-M, Ledru P (2009) A review of the pre-Permian geology of the Variscan French Massif Central. *CR Geosci* 341:202–213. <https://doi.org/10.1016/j.crte.2008.12.001>
- Gaudel T (2016) Conditions de pic de métamorphisme enregistrés par les gneiss de l'unité de la Dronne (Limousin): les limites de la fusion partielle dans le l'Ouest du Massif Central Français. Master's thesis, University of Orléans, pp 28
- Gébelin A, Brunel M, Monié P, Faure M, Arnaud N (2007) Transpressional tectonics and Carboniferous magmatism in the Limousin, Massif Central, France: Structural and 40Ar/39Ar investigations. *Tectonics* 26:1–27. <https://doi.org/10.1029/2005TC001822>
- Goodenough KM, Lusty PAJ, Roberts NMW, Key RM, Garba A (2014) Post-collisional Pan-African granitoids and rare metal pegmatites in western Nigeria: age, petrogenesis, and the “pegmatite

- conundrum." *Lithos* 200–201:22–34. <https://doi.org/10.1016/j.lithos.2014.04.006>
- Haapala I, Lukkari S (2005) Petrological and geochemical evolution of the Kymi stock, a topaz granite cupola within the Wiborg rapakivi batholith, Finland. *Lithos* 80:347–362
- Hammouda T, Pichavant M (1999) Kinetics of melting of fluorophlogopite-quartz pairs at 1 atmosphere. *Eur J Mineral* 11:637–653
- Hammouda T, Pichavant M (2000) Melting of fluorophlogopite-plagioclase pairs at 1 atmosphere. *Eur J Mineral* 12:315–328
- Hanson GN (1978) The application of trace elements to the petrogenesis of igneous rocks of granitic composition. *Earth Planet Sci Lett* 38:26–43
- Harlaux M, Mercadier J, Marignac C, Peiffert C, Cloquet C, Cuney M (2018) Tracing metal sources in peribatholithic hydrothermal W deposits based on the chemical composition of wolframite: the example of the Variscan French Massif Central. *Chem Geol* 479:58–85. <https://doi.org/10.1016/j.chemgeo.2017.12.029>
- Harris NBW, Inger S (1992) Trace element modelling of pelite-derived granites. *Contrib Mineral Petrol* 110:46–56
- Harris NBW, Ayres M, Massey J (1995) Geochemistry of granitic melts produced during the incongruent melting of muscovite: implications for the extraction of Himalayan leucogranite magmas. *J Geophys Res* 100(B8):14767–15777
- Hertogen J, Gijbels R (1976) Calculation of trace element fractionation during partial melting. *Geochim Cosmochim Acta* 40:313–322
- Holyoke CW, Rushmer T (2002) An experimental study of grain scale melt segregation mechanisms in two common crustal rock types. *J Metamorph Geol* 20:493–512
- Huang WL, Wyllie PJ (1974) Melting relations of muscovite with quartz and sanidine in the $K_2O-Al_2O_3-SiO_2-H_2O$ system to 30 kilobars and an outline of paragonite melting relations. *Am J Sci* 274:378–395
- Hulsbosch N, Hertogen J, Dewaele S, Andre L, Muchez P (2014) Alkali metal and rare earth element evolution of rock-forming minerals from the Gatumba area pegmatites (Rwanda): quantitative assessment of crystal-melt fractionation in the regional zonation of pegmatite groups. *Geochim Cosmochim Acta* 132:349–374
- Hulsbosch N (2019) Nb-Ta-Sn-W Distribution in Granite-related Ore Systems: fractionation mechanisms and examples from the Karagwe-Ankole belt of Central Africa. In *Ore Deposits American Geophysical Union (AGU)*, pp 75–107
- Icenhower J, London D (1995) An experimental study of element partitioning among biotite, muscovite, and coexisting peraluminous silicic melt at 200 MPa (H_2O). *Am Miner* 80:1229–1251
- Icenhower J, London D (1996) Experimental partitioning of Rb, Cs, Sr, and Ba between alkali feldspar and peraluminous melt. *Am Miner* 81:719–734
- Jahns RH, Burnham CW (1969) Experimental studies of pegmatite genesis; I, A model for the derivation and crystallization of granitic pegmatites. *Econ Geol* 64:843–864
- Johannes W, Holtz F (1996) Petrogenesis and experimental petrology of granitic rocks. In: Wyllie PJ, El Goresy A, von Engelhardt W, Hahn T (eds) *Minerals and rocks 22*. Springer-Verlag, Berlin, p 335
- Kesraoui M, Marignac C, Verkaeren J (2000) L'évolution tardi-magmatique des granites à métaux rares: l'exemple de la coupole de Tin-Amzi (Hoggar-Algérie). *Bulletin De La Société Géologique De L'algerie* 11(2):195–216
- Kinnaid J, Bowden P, Ixer RA, Odling NWA (1985) Mineralogy, geochemistry and mineralization of the Ririwai complex, northern Nigeria. *J Afr Earth Sc* 3:185–222
- Kontak DJ (1990) The East Kemptville topaz-muscovite leucogranite, Nova Scotia; I. Geological setting and whole-rock geochemistry. *Can Mineral* 28(4):787–825
- Konzett J, Hauzenberger C, Ludwig T, Stalder R (2018) Anatectic granitic pegmatites from the eastern Alps: a case study of variable rare metal enrichment during high-grade regional metamorphism. II. Pegmatite staurolite as an indicator of anatectic pegmatite parent melt formation—a field and experiment study. *Can Mineral* 56:603–624
- Kovalenko VI, Tsaryeva GM, Goreglyad AV, Yarmolyuk VV, Troitsky VA, Hervig RL, Farmer GL (1995) The peralkaline granite-related Khaldzan-Buregtey rare metal (Zr, Nb, REE) deposit, western Mongolia. *Econ Geol* 90:530–547
- Kovalenko V I, Kovalenko N I (1976) Ongonites (topaz bearing quartz keratophyre) – subvolcanic analogues of rare metal Li-F granites: Nauka, Moskva, pp 124 (in Russian)
- Lardeaux J-M (2014) Deciphering orogeny: a metamorphic perspective. Examples from European Alpine and Variscan belts. Part II: Variscan metamorphism in the French Massif Central—a review. *Bull Soc Géol France* 185(5):281–310
- Le Breton N, Thompson AB (1988) Fluid-absent (dehydration) melting of biotite in metapelites in the early stage of crustal anatexis. *Contrib Miner Petrol* 99:226–237
- Le Fort P, Cuney M, Deniel C, France-Lanord C, Sheppard SMF, Upreti BN, Vidal P (1987) Crustal generation of the Himalayan leucogranites. *Tectonophysics* 134:39–57
- Linnen RL, Pichavant M, Holtz F, Burgess S (1995) The effect of f_{O_2} on the solubility, diffusion, and speciation of tin in haplogranitic melt at 850 °C and 2 kbar. *Geochim Cosmochim Acta* 59:1579–1588
- Linnen R L, Cuney M (2005) Granite-related rare-element deposits and experimental constraints on Ta-Nb-W-Sn-Zr-Hf mineralization. In: Linnen RL and Samson IM (eds) *Rare-element geochemistry and mineral deposits*. In: Geological Association of Canada, GAC, Short Course Canada
- London D (2005) Granitic pegmatites: an assessment of current concepts and directions for the future. *Lithos* 80:281–303. <https://doi.org/10.1016/j.lithos.2004.02.009>
- London D (2008) Pegmatites. *Canadian Mineralogist Special Publication*, 10, pp 368
- Lotout C, Pitra P, Poujol M, Van Den Driessche J (2017) Ordovician magmatism in the Lévézou massif (French Massif Central): tectonic and geodynamic implications. *Int J Earth Sci* 106:501–515
- Luth WC (1969) The system $NaAlSi_3O_8-SiO_2$ and $KAlSi_3O_8$ to 20kb and the relationship between H_2O content p_{H_2O} and p_{total} in granitic magmas. *Am J Sci* 267A:325–341
- Manning DAC, Hill PI (1990) The petrogenetic and metallogenetic significance of topaz granite from the Southwest England orefield. *Geol Soc Am Spec Pap* 246:51–69
- Mayne MJ, Moyen J-F, Stevens G, Kaisleniemi L (2016) Rcrust: a tool for calculating path-dependent open system processes and application to melt loss. *J Metamorph Geol* 34:663–682
- Melton J, Faure M, Cocherie A (2009) Monazite U-Th/Pb chemical dating of the Early Carboniferous syn-kinematic MP/MT metamorphism in the Variscan French Massif Central. *Bull Soc Géol France* 180(3):283–292
- Melton J, Gloaguen E, Frei D (2015) Rare elements (Li-Be-Ta-Sn-Nb) magmatism in the European Variscan belt, a review. *SGA, Ressources minérales dans un monde durable*, Nancy, France pp 807–810
- Michaud JA-S, Pichavant M (2020) Magmatic fractionation and the magmatic-hydrothermal transition in rare metal granites: evidence from Argemela (Central Portugal). *Geochim Cosmochim Acta* 289:130–157. <https://doi.org/10.1016/j.gca.2020.08.022>
- Michaud JA-S (2019) Rare metal granites: origin, emplacement and mechanisms of the magmatic-hydrothermal transition. Insights from the Argemela rare metal granite (Portugal) and an experimental study. PhD thesis, University of Orléans, pp 365

- Misra S, Burlini L, Burg J-P (2009) Strain localization and melt segregation in deforming metapelites. *Phys Earth Planet Inter* 177:173–179. <https://doi.org/10.1016/j.pepi.2009.08.011>
- Montel J-M, Vielzeuf D (1994) Partial melting of metagreywackes 1. Fluid absent experiments and phase relationships. *Contrib Miner Petrol* 117:375–393
- Montel J-M, Vielzeuf D (1997) Partial melting of metagreywacke-2: compositions of minerals and melts. *Contrib Miner Petrol* 128:176–196
- Montel J-M, Weber C, Barbey P, Pichavant M (1986) Thermo-barométrie du domaine anatectique du Velay (Massif Central, France) et conditions de genèse des granites tardi-migmatitiques. *Comptes Rendus Académie Des Sciences Paris* 302:647–652
- Muller A, Romer R, Pedersen R-B (2017) The Sveconorwegian pegmatite province—thousands of pegmatites without parental granites. *Can Mineral* 55:283–315
- Nabelek PI (1999) Trace element distribution among rock-forming minerals in Black Hills migmatites, South Dakota: a case for solid-state equilibrium. *Am Miner* 84:1256–1269
- Nabelek PI, Russ-Nabelek C, Denison JR (1992a) The generation and crystallization conditions of the Proterozoic Harney Peak Leucogranite, Black Hills, South Dakota, USA: Petrologic and geochemical constraints. *Contrib Mineral Petrol* 110:173–191
- Nabelek PI, Russ-Nabelek C, Haeussler GT (1992b) Stable isotope evidence for the petrogenesis and fluid evolution in the Proterozoic Harney Peak leucogranite, Black Hills, South Dakota. *Geochim Cosmochim Acta* 56:403–417
- Neiva ANAMR (2002) Portuguese granites associated with Sn-W and Au mineralizations. *Bull Geol Soc Finl* 79(1–2):79–101
- PatíñoDouce AE, Harris N (1998) Experimental constraints on Himalayan anatexis. *J Petrol* 39(4):689–710
- Pearce NJG, Perkins WT, Westgate JA, Gorton MP, Jackson SE, Neal CR, Chenery SP (1997) A compilation of new and published major and trace element data for NIST SRM 610 and NIST SRM 612 glass reference materials. *Geostand News* 21(1):115–144
- Pichavant M, Valencia Herrera J, Boulmier S, Briquieu L, Joron JL, Juteau M, Marin L, Michard A, Sheppard SMF, Treuil M, Vernet M (1987) The Macusani glasses, SE Peru: evidence of chemical fractionation in peraluminous magmas. In *Magmatic processes: physicochemical principles* (ed. B.O. Mysen). *Geochem Soc Spec Public* 1:359–373
- Pichavant M, Kontak DJ, Herrera V, Clark AH (1988a) The Miocene-Pliocene Macusani Volcanics, SE Peru. I. Mineralogy and magmatic evolution of a two-mica aluminosilicate-bearing ignimbrite suite. *Contrib Miner Petrol* 100:300–324
- Pichavant M, Kontak D, Briquieu L, Valencia Herrera J, Clark AH (1988b) The Macusani Volcanics, SE Peru: II. Geochemistry and origin of a felsic peraluminous magma. *Contrib Miner Petrol* 100:325–338
- Pichavant M, Villaros A, Deveaud S, Scaillet B, Lahfafi M (2016) The influence of redox state on mica crystallization in leucogranite and pegmatitic liquids. *Can Mineral* 54:559–581
- Pillet D, Chevenoy M, Bélanger M (1992) Pétrologie du granite peralcalin du lac Brisson, Labrador central, Nouveau-Québec. 1. Mode de mise en place et évolution chimique. *Revue Canadienne Des Sciences De La Terre* 29:353–372
- Raimbault L (1987) Genèse des granites à métaux rares : Revue comparative des modèles géochimiques. *Géol Fr* 3:101–108
- Raimbault L, Burnol L (1998) The Richemont rhyolite dyke, Massif Central, France; a subvolcanic equivalent of rare-metal granites. *Can Mineral* 36:265–282
- Raimbault L, Cuney M, Azencott C, Duthou J-L, Joron J (1995) Geochemical evidence for a multicistage magmatic genesis of Ta-Sn-Li mineralization in the granite at Beauvoir, French Massif Central. *Econ Geol* 90:548–576
- Renno A (1997) Zur Petrogenese der Albitgranite von Abu Dabbab und Nuweibi, Central Eastern Desert, Ägypten. Unpublished PhD Thesis, Technische Universität Berlin, Berlin, pp 216
- Robie RA, Hemingway BS, Fisher JR (1979) Thermodynamic properties of minerals and related substances at 298.15 K and 1 Bar (10^5 Pascals) pressure and at higher temperatures. *US Geological Survey Bulletin* 1452
- Roda E, Pesquera A, Velasco F, Fontan F (1999) The granitic pegmatites of the Fregeneda area (Salamanca, Spain): characteristics and petrogenesis. *Mineral Mag* 63:535–556
- Roda-Robles E, Villaseca C, Pesquera A, Gil-Crespo PP, Vieira R, Lima A, Garate-Olave I (2018) Petrogenetic relationships between Variscan granitoids and Li-(F-P)-rich aplite-pegmatites in the Central Iberian Zone: geological and geochemical constraints and implications for other regions from the European Variscides. *Ore Geol Rev* 95:408–430
- Romer R, Kroner U (2015) Sediment and weathering control on the distribution of Paleozoic magmatic tin-tungsten mineralization. *Miner Deposita* 50:327–338
- Rudnick RL, Gao S (2003) Composition of the continental crust. In: Rudnick RL (ed) *The crust*. Elsevier, Amsterdam, pp 1–70
- Rushmer T (2001) Volume change during partial melting reactions: implications for melt extraction, melt geochemistry and crustal rheology. *Tectonophysics* 342:389–405
- Scaillet B, Pichavant M, Roux J (1995) Experimental crystallization of leucogranite magmas. *J Petrol* 36:663–705
- Schmitt AK, Emmermann R, Trumbull RB, Bühn B (2000) Petrogenetic aspects of metaluminous and per-alkaline granites in the Brandberg anorogenic complex, Namibia: evidence for major mantle contribution. *J Petrol* 41:1207–1239
- Schneider CA, Rasband WS, Eliceiri KW (2012) NIH image to ImageJ: 25 years of image analysis. *Nat Methods* 9:671–675
- Shaw DM (1970) Trace element fractionation during anatexis. *Geochim Cosmochim Acta* 34:237–243
- Shaw RA, Goodenough KM, Roberts NMW, Horstwood MSA, Chenery SR, Gunn AG (2016) Petrogenesis of rare-metal pegmatites in high-grade metamorphic terranes: a. *Precamb Res* 281:338–362. <https://doi.org/10.1016/j.precamres.2016.06.008>
- Shearer CK, Papike JJ, Laul JC (1987) Mineralogical and chemical evolution of a rare-element granite-pegmatite system: Harney Peak granite, Black Hills, south Dakota. *Geochim Cosmochim Acta* 51:473–486
- Shearer CK, Papike JJ, Jolliff BL (1992) Petrogenetic links among granites and pegmatites in the Harney Peak rare-element granite-pegmatite system, Black Hills, south Dakota. *Can Mineral* 30:785–809
- Simons B, Shail RK, Andersen JCØ (2016) The petrogenesis of the Early Permian Variscan granites of the Cornubian Batholith: lower plate post-collisional peraluminous magmatism in the Rhe-nohercynian Zone of SW England. *Lithos* 260:76–94. <https://doi.org/10.1016/j.lithos.2016.05.010>
- Simons B, Andersen JCØ, Shail RK, Jenner FE (2017) Fractionation of Li, Be, Ga, Nb, Ta, In, Sn, Sb, W and Bi in the peraluminous Early Permian Variscan granites of the Cornubian Batholith: precursor processes to magmatic-hydrothermal mineralisation. *Lithos* 278–281:491–512. <https://doi.org/10.1016/j.lithos.2017.02.007>
- Spear FS (1995) *Metamorphic phase equilibria and pressure-temperature-time paths*, 2nd edn. Mineralogical Society of America, Washington
- Stepanov AS, Hermann J (2013) Fractionation of Nb and Ta by biotite and phengite: implications for the “missing Nb paradox.” *Geology* 41:303–306
- Stepanov A, Mavrogenes JA, Meffre S, Davidson P (2014) The key role of mica during igneous concentration of tantalum. *Contrib Mineral Petrol* 167:1009. <https://doi.org/10.1007/s00410-014-1009-3>

- Stevens G, Villaros A, Moyer J-F (2007) Selective peritectic garnet entrainment as the origin of geochemical diversity in S-type granites. *Geology* 35:9–12. <https://doi.org/10.1130/G22959A.1>
- Stewart DB (1978) Petrogenesis of lithium-rich pegmatites. *Am Miner* 63:970–980
- Taylor JR, Wall VJ, Pownceby MI (1992) The calibration and application of accurate redox sensors. *Am Miner* 77:284–295
- Thompson AB (1982) Dehydration Melting of Pelitic rocks and the generation of H₂O-undersaturated granitic liquids. *Am J Sci* 282:1567–1595
- Tumarkina E, Misra S, Burlini L, Connolly JAD (2011) An experimental study of the role of shear deformation on partial melting of a synthetic metapelite. *Tectonophysics* 503:92–99
- Turpin L, Leroy JL, Sheppard SMF (1990) Isotopic (O, H, C, Sr, Nd) of superimposed barren and U-bearing hydrothermal systems in a Hercynian granite, Massif Central, France. *Chem Geol* 88:85–98
- Vielzeuf D, Clemens JD (1992) The fluid-absent melting of phlogopite + quartz: experiments and models. *Am Miner* 77:1206–1222
- Vielzeuf D, Holloway JR (1988) Experimental determination of the fluid-absent melting relations in the pelitic system: consequences for crustal differentiation. *Contrib Mineral Petrol* 98:257–276
- Vielzeuf D, Montel J-M (1994) Partial melting of metagreywackes. Part I. Fluid-absent experiments and phase relationships. *Contrib Mineral Petrol* 177:375–393
- Vielzeuf D, Schmidt MW (2001) Melting relations in hydrous systems revisited: application to metapelites, metagreywackes and metabasalts. *Contrib Mineral Petrol* 141:251–267
- Villaros A, Pichavant M (2019) Mica-liquid trace elements partitioning and the granite-pegmatite connection: the St-Sylvestre complex (Western French Massif Central). *Chem Geol* 528:119265
- Villaros A, Laurent O, Couzinié S, Moyer J-F, Mintrone M (2018a) Plutons and domes: the consequences of anatectic magma extraction—example from the southeastern French Massif Central. *Int J Earth Sci* 107:2819–2842. <https://doi.org/10.1007/s00531-018-1630-x>
- Villaros A, Pichavant M, Michaud J (2018b) Experimental melting of ortho- and paragneiss under fluid-present conditions: melt compositions (major and trace elements) and implications for granite genesis. EMPG, Clermont-Ferrand, p 130
- Wolf M, Romer RL, Franz L, López-Moro FJ (2018) Tin in granitic melts: the role of melting temperature and protolith composition. *Lithos* 310–311:20–30
- Wolf M (2018) The role of partial melting on trace element and isotope systematics of granitic melts. Potsdam, PhD p 139
- Yakymchuk C, Brown M (2014) Behaviour of zircon and monazite during crustal melting. *J Geol Soc Lond* 171:465–479
- Yang P, Rivers T (2000) Trace element partitioning between coexisting biotite and muscovite from metamorphic rocks, Western Labrador: structural, compositional and thermal controls. *Geochim Cosmochim Acta* 64:1451–1472
- Yardley BWD (2009) The role of water in the evolution of the continental crust. *J Geol Soc Lond* 166:585–600. <https://doi.org/10.1144/0016-76492008-101>
- Yuan S, Williams-Jones AE, Romer RL, Zhao P, Mao J (2019) Protolith-related thermal controls on the decoupling of Sn and W in Sn-W metallogenic provinces: insights from the Nanling region, China. *Econ Geol* 114(5):1005–1012
- Zhang Y, Ni H, Chen Y (2010) Diffusion data in silicate melts. *Rev Mineral Geochem* 72:311–408
- Zhu J-C, Li R-K, Li F-C, Xiong X-L, Zhou F-Y, Huang X-L (2001) Topaz-albite granites and rare-metal mineralization in the Limu District, Guangxi Province, southeast China. *Miner Deposita* 36(5):393–405

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.